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GenCore version 5.1.7

OM protein - protein search, using sw model

Run on: February 28, 2006, 15:24:01 ; Search time 39 Seconds

(without alignments)
407,071 Million cell updates/sec

Title: US-10-706-701-1

Sequence: 1 APPRLICDSRVIERYLLEAK.....SNFLRGKLUWGRACRTG 165

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 283416 seqs, 96216763 residues

Total number of hits satisfying chosen parameters: 283416

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

PIR-80:*

1: pir1:*

2: pir2:*

3: pir3:*

4: pir4:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match Length	DB ID	Description
1	846	100.0	193	ZUHU erythropoietin pre
2	764.5	90.4	192	JQ0173 erythropoietin pre
3	759.5	89.8	192	184133 erythropoietin pre
4	701	82.9	192	146083 erythropoietin pre
5	683.5	81.0	194	14601 erythropoietin pre
6	680.5	80.5	195	2 JC7699 erythropoietin pre
7	678	80.1	190	2 146578 erythropoietin - p
8	638	75.4	175	2 146199 erythropoietin - h
9	90	10.6	353	2 G02729 thrombopoietin pre
10	89	10.5	353	2 180105 erythropoietin pre
11	88	10.4	323	2 AB0323 ribonucleoside-diphosphate binding rec
12	87.5	10.3	346	2 AB0559 megakaryocyte growth
13	86	10.2	286	2 A55530 probable 2-hydroxy
14	83	9.8	296	2 A10443 thrombopoietin - h
15	83	9.8	339	2 AB3274 UPF-N-acetylpyruvo
16	80.5	9.5	3033	1 GNWYU8 genome polyprotein
17	79.5	9.4	1829	1 P0581 probable sensory h
18	79	9.2	480	2 S56339 ribosomal protein
19	79	9.3	813	2 AF0526 ATP-dependent heli
20	78.5	9.3	897	2 A54696 EGFR receptor sub
21	78.5	9.3	92	2 T3450 ABC transporter AT
22	78	9.2	348	2 conserved hypothetical protein
23	78	9.2	455	2 AG2919 methylamine utilize
24	78	9.2	455	2 H97693 probable copper-tr
25	78	9.2	477	1 S36741 hypothetical prote
26	77	9.2	242	2 AD1928 hypothetical prote
27	77	9.1	451	2 ST5569 bacterioferritin X
28	76.5	9.0	154	2 H82810

ALIGNMENTS

RESULT 1
ZUHU erythropoietin precursor [validated] - human
C;Species: Homo sapiens (man)
C;Date: 27-Nov-1985 #Sequence revision 27-Nov-1985 #text_change 09-Jul-2004
C;Accession: A01855; A24744; A25384; A22120; S56178
R;Jacobs, K.; Shoemaker, C.; Rutherford, R.; Neill, S.D.; Kaufman, R.J.; Mufson, A.; Se
Nature 313, 806-810, 1985
A;Title: Isolation and characterization of genomic and cDNA clones of human erythropoietin gene.
A;Reference number: A01855; MUID:8513789; PMID:3038366
A;Accession: A01855
A;Molecule type: mRNA; DNA
A;Residues: 1-193 <JAC>
A;Cross-references: UNIPROT:P01588; UNIPARC:UPI000033477; GB:X02157; GB:X02158
R;Lin, F.K.; Suggs, S.; Lin, C.H.; Browne, J.K.; Smalling, R.; Egrie, J.C.; Chen, K.K.; Se
Proc. Natl. Acad. Sci. U.S.A. 82, 7580-7584, 1985
A;Title: Cloning and expression of the human erythropoietin gene.
A;Reference number: A24744; MUID:86067948; PMID:3065178
A;Accession: A24744
A;Molecule type: DNA
A;Residues: 1-193 <LN>
A;Cross-references: UNIPARC:UPI000033477; GB:M11319; NID:9182197; PIDN:AAA52400_1; PID
R;Lai, P.H.; Everett, R.; Wang, P.F.; Arakawa, T.; Goldwasser, E.
J. Biol. Chem. 261, 3116-3121, 1986
A;Title: Structural characterization of human erythropoietin.
A;Reference number: A25384; MUID:86140080; PMID:3949763
A;Molecule type: protein
A;Residues: 28-86; Q; 87-193 <IAI>
A;Cross-references: UNIPARC:UPI00001736A2
A;Experimental source: urine
A;Note: forms with the carboxyl-terminal residue and the four carboxyl-terminal resi
R;Yanagawa, S.; Hirade, K.; Ohnoto, H.; Sasaki, R.; Chiba, H.; Ueda, M.; Goto, M.
J. Biol. Chem. 259, 2707-2710, 1984
A;Title: Isolation of human erythropoietin with monoclonal antibodies.
A;Reference number: A22210; MUID:84135751; PMID:6698989
A;Accession: A22210
A;Molecule type: protein
A;Residues: 28-29; X; 31-33; 'L'; 35-50; 'X'; 52-53; 'D'; 55; 'G'; 57 <YAN>
A;Cross-references: UNIPARC:UPI0000142781
R;Matsumoto, S.; Ikura, K.; Ueda, M.; Sasaki, R.
Plant Mol. Biol. 27, 1163-1172, 1995
A;Title: Characterization of a human glycoprotein (erythropoietin) produced in cultured
A;Reference number: S56178; MUID:952884365; PMID:7766897
A;Accession: S56178
A;Molecule type: protein
A;Residues: 28-33; X; 35-37 <WMS>
A;Cross-references: UNIPARC:UPI00001736A3
C;Comment: Erythropoietin is produced by kidney or liver of adult mammals and by liver
C;Genetics:
A;Gene: GDB:EPO
A;Cross-references: GDB:119110; OMM:133170

RESULT 3
 A;Map position: 7q21.3-7q22.1
 A;Introns: 5/1; 53/3; 82/3; 142/3
 C;Function:
 C;Description: the primary inducer of erythrocyte formation
 C;Superfamily: erythropoietin
 C;Keywords: erythropoietin; glycoprotein; hormone; kidney; liver
 F;1-27/Domain: signal sequence #status predicted <SIG>
 F;28-193/Product: erythropoietin #status experimental <MAT>
 F;51-65,110/Binding site: carbohydrate (Asn) (covalent) #status experimental
 F;153/Binding site: carbohydrate (Ser) (covalent) #status experimental

Query Match 100.0%; Score 846; DB 1; Length 193;
 Best Local Similarity 100.0%; Pred. No. 8,6e-74; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSVRVLRVYLREAKAENITTCGAHCNSLNENITVPDKTKNFYAWKRMENVGQAA 60
 Db 28 APPRLICDSVRVLRVYLREAKAENITTCGAHCNSLNENITVPDKTKNFYAWKRMENVGQAA 87

Qy 61 VEWMOGLALISSEAVLVRGQALLVNSQPPWEPLOLHQDVKAVSGLSRSLTLLRAIGAQEAIIS 120
 Db 88 VEWMOGLALISSEAVLVRGQALLVNSQPPWEPLOLHQDVKAVSGLSRSLTLLRAIGAQEAIIS 147

Qy 121 PPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 165
 Db 148 PPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 192

RESULT 2
 JQ0173 erythropoietin precursor - crab-eating macaque
 C;Species: Macaca fascicularis (crab-eating macaque)
 C;Date: 07-Sep-1990 #sequence_revision 15-Nov-1996 #text_change 09-Jul-2004
 C;Accession: JQ0173
 R;Jin, F.K.; Lin, C.H.; Lai, P.H.; Browne, J.K.; Egrie, J.C.; Smalling, R.; Fox, G.M.; C;Title: Monkey erythropoietin gene: cloning, expression and comparison with the human gene 44, 201-209, 1986
 A;Reference number: JQ0173; MUID:87055236; PMID:2877922
 A;Accession: JQ0173
 A;Molecule type: mRNA
 A;Residues: 1-192 <LIN>
 A;Cross-references: UNIPROT:P07865; UNIPARC:UPI000012A0B2; GB:MI18189; GB:MI5818; GB:MI58
 A;Experimental source: kidney
 C;Comment: This protein is the principal hormone involved in the regulation of erythrocyte formation
 C;Function: the primary inducer of erythrocyte formation
 C;Keywords: erythropoietin
 F;1-27/domain: signal sequence #status predicted <SIG>
 F;28-192/Product: erythropoietin #status predicted <MAT>
 F;34-187,56-60/Disulfide bonds: #status predicted
 F;51,65,110/Binding site: carbohydrate (Asn) (covalent) #status predicted
 F;152/Binding site: carbohydrate (Ser) (covalent) #status predicted

Query Match 89.8%; Score 759.5; DB 1; Length 192;
 Best Local Similarity 90.3%; Pred. No. 1,7e-65; Indels 6; Gaps 1; Mismatches 149; Conservative 9; Mismatches 9; Indels 1; Gaps 1;

Qy 1 APPRLICDSVRVLRVYLREAKAENITTCGAHCNSLNENITVPDKTKNFYAWKRMENVGQAA 60
 Db 28 APPRLICDSVRVLRVYLREAKAENITTCGAHCNSLNENITVPDKTKNFYAWKRMENVGQAA 87

Qy 61 VEWMOGLALISSEAVLVRGQALLVNSQPPWEPLOLHQDVKAVSGLSRSLTLLRAIGAQEAIIS 120
 Db 88 VEWMOGLALISSEAVLVRGQALLVNSQPPWEPLOLHQDVKAVSGLSRSLTLLRAIGAQ-EAIIS 147

Qy 121 PPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 165
 Db 147 PPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 191

RESULT 4
 I46083 erythropoietin precursor - cat (fragment)
 C;Species: Felis silvestris catus (domestic cat)
 C;Date: 16-Aug-1996 #sequence_revision 15-Nov-1996 #text_change 09-Jul-2004
 C;Accession: I46083
 R;Men, D.; Boissel, J.
 Blood 82, 1507-1516, 1993
 A;Title: Erythropoietin structure-function relationships: High degree of sequence homology
 A;Reference number: I46083; MUID:93372347; PMID:8364201
 A;Accession: I46083
 A;Status: translated from GB/EMBL/DDJB
 A;Molecule type: mRNA
 A;Residues: 1-188 <WEN>
 A;Cross-references: UNIPROT:P33708; UNIPARC:UPI00016C43A; GB:II10606; NID:9163820; PIDN
 C;Comment: Erythropoietin is produced by kidney or liver of adult mammals and by liver
 C;Function:
 C;Description: the primary inducer of erythrocyte formation
 C;Superfamily: erythropoietin
 C;Keywords: erythropoiesis; glycoprotein; hormone; kidney; liver
 F;1-22/domain: signal sequence (fragment) #status predicted <SIG>
 F;23-188/Product: erythropoietin #status predicted <MAT>
 F;28-183,51-55/Disulfide bonds: #status predicted
 F;46,60,105/Binding site: carbohydrate (Asn) (covalent) #status predicted
 F;148/Binding site: carbohydrate (Ser) (covalent) #status predicted

Query Match 84.3%; Score 713; DB 1; Length 188;
 Best Local Similarity 84.2%; Pred. No. 4,9e-61; Indels 0; Gaps 0;
 Matches 139; Conservative 9; Mismatches 17; Indels 0; Gaps 0;

Qy 121 PPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 165
 Db 147 LPDASAAPIRTTADTPRLFRVYSNPLRGKLYGEACRGD 191

QY 1 APPRLICDSRVLERYLRAKEAENITGCAHCISLNENITPDTKVNFIYAWKRMVEQQA 60
 Db 23 APPRLICDSRVLERYLRAKEAENITGCAHCISLNENITPDTKVNFIYAWKRMVEQQA 82
 QY 61 VEWQOGLALISEAVLRGQLVNSQPWEPLQHDKAVSGIRSLSLTLLRGAKEATS 120
 Db 83 VEWQOGLALISEAVLRGQLVNSQPWEPLQHDKAVSGIRSLSLTLLRGAKEATS 142
 QY 121 PPDAA-SAAPLRTTADFRKLFRVYSNFLRGKLYTGACRTGD 165
 Db 143 LPPDATSAAPLRTTADFRKLFRVYSNFLRGKLYTGACRTGD 187

RESULT 5

S28148 erythropoietin precursor - rat

C;Species: *Rattus norvegicus* (Norway rat)

C;Date: 22-Nov-1993 #sequence_revision 15-Nov-1996 #text_change 09-Jul-2004

C;Accession: S28148; I6743

R;Nagao, M.; Suga, H.; Okano, M.; Masuda, S.; Narita, H.; Ikura, K.; Sasaki, R.

Blood, 82, 1507-1516, 1993

C;Title: Nucleotide sequence of rat erythropoietin.

A;Reference number: S28148; MUID:93042015; PMID:1420369

A;Molecule type: mRNA

A;Residues: 1-192 <WAG>

A;Cross-references: UNIPROT:P29676; UNIPARC:UPI000012A0B5; GB:D10763; NID:9220735; PIDN:

R;Wen, D.; Boissel, J.

Blood, 82, 1507-1516, 1993

A;Accession: I62743

A;Reference number: I46033; MUID:93372347; PMID:8364201

A;Molecule type: mRNA

A;Residues: 4-92 <RES>

A;Cross-references: UNIPARC:UPI0000170949; GB:L10508; NID:9204060; PIDN:AAA41126.1; PID:

C;Function: the primary inducer of erythrocyte formation

A;Description: the primary inducer of erythrocyte formation

C;Superfamily: erythropoietin

C;Keywords: erythropoiesis; glycoprotein; hormone; kidney; liver

F;1-27//Domain: Signal sequence #status predicted <SIG>

F;28-194//Product: erythropoietin #status predicted <MAT>

F;34-189, 56-60//Disulfide bonds: #status predicted

F;51, 65, 110//Binding site: carbohydrate (Asn) (covalent) #status predicted

F;1-54//Binding site: carbohydrate (Ser) (covalent) #status predicted

C;Comment: Erythropoietin is produced by kidney or liver of adult mammals and by liver of

C;Function:

A;Description: the primary inducer of erythrocyte formation

C;Superfamily: erythropoietin

C;Keywords: erythropoiesis; glycoprotein; hormone; kidney; liver

F;1-16//Domain: signal sequence #status predicted <SIG>

F;33-187, 55-165//Disulfide bonds: #status predicted (Asn) (covalent) #status predicted

F;50, 64, 109//Binding site: carbohydrate

Query Match 82.9% Score 701; DB 1; Length 192; Matches 136; Conservative 82.4%; Pred. No. 7, 1e-50; Mismatches 13; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLRAKEAENITGCAHCISLNENITPDTKVNFIYAWKRMVEQQA 60
 Db 27 APPRLICDSRVLERYLRAKEAENITGCAHCISLNENITPDTKVNFIYAWKRMVEQQA 86
 QY 61 VEWQOGLALISEAVLRGQLVNSQPWEPLQHDKAVSGIRSLSLTLLRGAKEATS 120
 Db 148 LPPDATSAAPLRTTADFRKLFRVYSNFLRGKLYTGACRTGD 193

RESULT 7

A24902 erythropoietin precursor - mouse

C;Species: *Mus musculus* (house mouse)

C;Date: 25-Oct-1987 #sequence_revision 15-Nov-1996 #text_change 09-Jul-2004

C;Accession: A24902; A24901

R;Shoemaker, C.B.; Mitsock, L.D.

Nol. Cell. Biol. 6, 849-858, 1986

A;Title: Murine erythropoietin gene: cloning, expression, and human gene homology.

A;Reference number: A24902; MUID:87039105; PMID:3773894

A;Accession: A24902

A;Molecule type: DNA

A;Residues: 1-192 <SHO>

A;Cross-references: UNIPROT:P07321; UNIPARC:UPI00001736A4

A;Note: the authors translated the codon TTA for residue 12 as Phe, TTA for residue 43 as Leu, TTA for residue 146 as Ile.

R;McDonald, J.D.; Lin, F.K.; Goldwasser, E.

Nol. Cell. Biol. 6, 842-848, 1986

A;Title: Cloning, sequencing, and evolutionary analysis of the mouse erythropoietin gene

A;Reference number: A24901; MUID:87039104; PMID:3022133

A;Accession: A24901

A;Molecule type: DNA

A;Residues: 1-67, 'P', 69-192 <MD>

A;Cross-references: UNIPARC:UPI000029308; GB:M12930; NID:9193086; PIDN:AAA3750.1; PID:

C;Species: *Ovis orientalis aries*, Ovis ammon aries (domestic sheep)

C;Date: 16-Aug-1996 #sequence_revision 15-Nov-1996 #text_change 09-Jul-2004

C;Accession: I46401; I47077

R;Fu, P.; Evans, B.; Lim, G.B.; Moritz, K.; Wintour, E.M.

Mol. Cell. Endocrinol. 93, 107-116, 1993

A;Title: The sheep erythropoietin gene: molecular cloning and effect of hemorrhage on pl

A;Reference number: I46401; MUID:93351736; PMID:8349021

P;27-192/Product: erythropoietin #status predicted <MAT>
 P;33-187,55-165/disulfide bonds: #status predicted (covalent) #status predicted
 Query Match Similarity 80.5%; Score 681; DB 1; Length 192;
 Best Local Similarity 79.4%; Pred. No. 5..58; Mismatches 20; Indels 0; Gaps 0;
 Matches 131; Conservative 14; Mismatches 20; Indels 0; Gaps 0;

Query Match Similarity 80.1%; Score 678; DB 2; Length 190;
 Best Local Similarity 82.0%; Pred. No. 1..1e-57; Mismatches 21; Indels 2; Gaps 1;
 Matches 137; Conservative 7; Mismatches 21; Indels 2; Gaps 1;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 60
 Db 27 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 86

Qy 61 VEWMOGLALISEAVTQLRQALLNSSQPEPLQHDAVSGSLRSLTTLRALGAQKAIS 120
 87 IEWMOGLALISEATLQOQALLNSSQPEPLQHDAVSGSLRSLTSLRALGAQKAIP 146

Qy 121 PPDAASAPLRTTADTRKLFPRVYSNPLRGKLYTGEACRTGD 165
 Db 147 PPDTTPPAPLRTTADTRKLFPRVYANPLRGKLYTGEACRRD 191

RESULT 8
 JC7699
 erythropoietin - rabbit
 C;Species: Oryctolagus cuniculus (domestic rabbit)
 C;Date: 30-Sep-2001 #sequence_revision 30-Sep-2001 #text_change 22-Oct-2001
 C;Accession: JC7699
 R.Vilaita, A.; Wu, D.; Margalith, M.; Hobart, P.
 Biochem. Biophys. Res. Commun. 284, 823-827, 2001
 ATITLE: Rabbit EPO gene and cDNA: Expression of rabbit EPO after intramuscular injection
 AR: Reference number: JC7699; MUID:21280682; PMID:11396976
 A:Contents: Kidney
 A:Accession: JC7699
 A:Molecule type: DNA
 A:Residues: 1-195 <VIL>
 A:Cross-references: UNIPARC:UPI00008799F; GB:AF290943
 C:Comment: This protein, a heavily glycosylated 34k protein produced in the fetal liver
 C:Genetics:
 A:Gene: epo
 C:Superfamily: erythropoietin
 C:Keywords: glycoprotein; kidney

Query Match Similarity 80.4%; Score 680.5; DB 2; Length 195;
 Best Local Similarity 81.3%; Pred. No. 6..7e-58; Mismatches 18; Indels 1; Gaps 1;

Matches 135; Conservative 12; Mismatches 18; Indels 1; Gaps 1;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 60
 Db 29 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 88

Qy 61 VEWMOGLALISEAVTQLRQALLNSSQPEPLQHDAVSGSLRSLTTLRALGAQKAIS 120
 Db 89 VEWMOGLALISEAMLRQALLNSQPEPLQHDAVSGSLRSLTSLRALGAQKAIS 148

Qy 121 PPDAASAPLRTTADTRKLFPRVYSNPLRGKLYTGEACRTGD 165
 Db 149 PPEAKASSAALPLRTVAAADTICKLFPRVYANPLRGKLYTGEACRRD 194

RESULT 9
 I46578
 erythropoietin - pig (fragment)
 C;Species: Sus scrofa domestica (domestic pig)
 C;Date: 21-Feb-1997 #sequence_revision 21-Feb-1997 #text_change 09-Jul-2004
 C;Accession: I46578

R.Wen, D.; Boissel, J.
 Blood 82, 1507-1516, 1993

Query Match Similarity 75.4%; Score 638; DB 2; Length 175;
 Best Local Similarity 81.0%; Pred. No. 7..1e-54; Mismatches 16; Indels 0; Gaps 0;
 Matches 124; Conservative 13; Mismatches 16; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 60
 Db 23 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 82

Qy 61 VEWMOGLALISEAVTQLRQALLNSSQPEPLQHDAVSGSLRSLTTLRALGAQKAIS 120
 Db 83 LEVMOGLALISEATLQOQALLNSSQPEPLQHDAVSGSLRSLTSLRALGAQKAIS 142

Qy 121 PPDAASAPLRTTADTRKLFPRVYSNPLRGKLYTGEACRTGD 153
 Db 143 LPBEPASAPLRTVAAADTICKLFPRVYANPLRGKLYTGEACRRD 175

RESULT 10
 I46199
 erythropoietin - dog (fragment)
 C;Species: Canis lupus familiaris (dog)
 C;Date: 21-Feb-1997 #sequence_revision 21-Feb-1997 #text_change 09-Jul-2004
 C;Accession: I46199
 R.Wen, D.; Boissel, J.
 Blood, 82, 1507-1516, 1993
 ATITLE: Erythropoietin structure-function relationships: High degree of sequence homol.
 AR: Reference number: I46083; MUID:93372347; PMID:8364201
 A:Accession: I46199
 A:Status: preliminary; translated from GB/EMBL/DDBJ
 A:Molecule type: mRNA
 A:Residues: 1-175 <WEN>
 A:Cross-references: UNIPROT:P33707; UNIPARC:UPI00012A0B0; GB:L13027; NID:9290087; PIDN
 C;Cross-references: UNIPROT:P33707; UNIPARC:UPI00012A0B0; GB:L13027; NID:9290087; PIDN
 C;Superfamily: erythropoietin

Query Match Similarity 75.4%; Score 638; DB 2; Length 175;
 Best Local Similarity 81.0%; Pred. No. 7..1e-54; Mismatches 16; Indels 0; Gaps 0;
 Matches 124; Conservative 13; Mismatches 16; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 60
 Db 23 APPRLICDSRVLERYLLEAKEAENITTCGAEHCSSLNENITVPTDKVNFYAWKMEVQOA 82

Qy 61 VEWMOGLALISEAVTQLRQALLNSSQPEPLQHDAVSGSLRSLTTLRALGAQKAIS 120
 Db 83 LEVMOGLALISEATLQOQALLNSSQPEPLQHDAVSGSLRSLTSLRALGAQKAIS 142

Qy 121 PPDAASAPLRTTADTRKLFPRVYSNPLRGKLYTGEACRTGD 153
 Db 143 LPBEPASAPLRTVAAADTICKLFPRVYANPLRGKLYTGEACRRD 175

RESULT 11
 I46578
 erythropoietin - human
 C;Species: Homo sapiens (man)
 C;Date: 21-Dec-1996 #sequence_revision 06-Jun-1997 #text_change 05-Nov-1999
 C;Accession: G02729
 R.I'm, S.
 submitted to the EMBL Data Library, May 1996
 A:Reference number: H01637
 A:Accession: G02729
 A:Status: preliminary; translated from GB/EMBL/DDBJ
 A:Molecule type: mRNA
 A:Residues: 1-353 <IMX>
 A:Cross-references: UNIPARC:UPI00016B1CC; EMBL:U59493; NID:91401245; PIDN:AA03392.1;
 A:Gene: hTPO
 Query Match Score 90; DB 2; Length 353;

	Best local Similarity 26.3%; Pred. No. 0.67; Mismatches 75; Indels 20; Gaps 5; Matches 41; Conservative 20; Mismatches 75; Indels 20; Gaps 5;
QY	1 APRPLICDSRVRLVYLLEAKAENITGGAHSLNENITVNPDTKVNFTYAWKERNEMVGQA 60
Db	24 APP-ACDPLRISKLURSHVLSKLSQEVHLPTPLPVPVPSLGERMKOMEETKA 81
QY	61 VEWVOGLALLSEAVL-RGQALLVNSQWPFLQHVKAVSGRSLSLTLLRAIGAQKEA 118
Db	82 QDLIGAVTILLEGWMAARGQGLGPTCLSSINGQISEQVRULLGQLSQL---LGTQ--- 132
QY	119 ISPPDAASAPLRTTADTRKLUFVSYLERGKLK 154
Db	133 -LPPQG-----RTTAHKDPNAIFLSFOHLRGKVR 161
RESULT 12	
TR0105	thrombopoietin precursor - human N:Alternate names: c-Mpl ligand; megakaryocyte growth and development factor precursor
C:Species	Homo sapiens (man)
C:Date	24-May-1996 #sequence_revision 24-May-1996 #text_change 09-Jul-2004
C:Accession	I59201; I80105; S45331; S48707; I38672; I52610
R:Poster	D.C.; Sprecher, C.A.; Grant, P.J.; Kramer, J.M.; Kuijper, J.L.; Holly, R.D.; Proc. Natl. Acad. Sci. U.S.A. 91, 13023-13027, 1994
A:Title	Human thrombopoietin: gene structure, cDNA sequence, expression, and chromosomal localization
A:Reference	number: 159281; MUID:95108091; PMID:780916
A:Accession	I59281
A:Status	translated from GB/EMBL/DBJ
A:Molecule type	DNA
A:Residues	1-353 <RE>
A:Cross-references	UNIPARC:UPI00004A8D1; GB:I36052; PIDN: AAC37566.1; PIDN: A80225; UNIPROT:P40225; UNIPARC:UPI00004A8D1; GB:I36051; NID:9533214; PIDN: A80105; AB0001; MUID:95108091; PMID:780916
R:de Sauvage	F.J.; Habs, P.E.; Spencer, S.D.; Malloy, B.B.; Gurney, A.L.; Spencer, S.A.
D.V.	Baton, D.L.
Nature	369, 533-538, 1994
A:Title	Stimulation of megakaryocytopoiesis and thrombopoiesis by the c-Mpl ligand.
A:Reference	number: S45331; MUID:94261202; PMID:8202154
A:Accession	S45331
A:Status	preliminary
A:Molecule type	mRNA
A:Residues	1-353 <SAU>
A:Cross-references	UNIPARC:UPI00004A8D1; GB:I33410; NID:9506826; PIDN: AAA59857.1; PIDN: A80225; UNIPROT:Q8ZDC8; UNIPARC:UPI0000DCAA2; GB:AL590842; PIDN: CAC2889.1
R:Schma	Y.; Akabori, H.; Seki, N.; Hori, T.; Ogami, K.; Kato, T.; Shimada, Y.; Kawamura, F.
FBS: Lett.	33, 57-61, 1994
A:Title	Molecular cloning and chromosomal localization of the human thrombopoietin gene
A:Reference	number: S48740; MUID:95010765; PMID:7926023
A:Accession	S48740
A:Status	preliminary
A:Molecule type	DNA
A:Residues	1-353 <SOH>
A:Cross-references	UNIPARC:UPI00004A8D1; GB:D32046; NID:9577319; PIDN: BAA06807.1; PIDN: A80225; UNIPROT:Q8ZDC8; UNIPARC:UPI0000DCAA2; GB:AL590842; PIDN: CAC2889.1
R:Bartley	T.D.; Bogenberger, J.; Hunt, P.; Li, Y.S.; Lu, H.S.; Martin, F.; Chang, M.S.; Cell 77, 1117-1124, 1994
A:Title	Identification and cloning of a megakaryocyte growth and development factor that
A:Reference	number: A5463; MUID:94291201; PMID:8020099
A:Accession	I38672
A:Status	preliminary
A:Molecule type	mRNA
A:Residues	1-112, 'E', 114-353 <RE>
A:Cross-references	UNIPARC:UPI000016A0D7; EMBL:U11025; NID:9511223; PIDN:AAA50553.1; PIDN: A80225; UNIPROT:Q8ZDC8; UNIPARC:UPI00004A8D1; GB:D32046; NID:9577319; PIDN: BAA06807.1; PIDN: A80225; UNIPROT:Q8ZDC8; UNIPARC:UPI0000DCAA2; GB:AL590842; PIDN: CAC2889.1
R:Gurney	A.L.; Kiang, W.J.; Xie, M.H.; Maloy, B.E.; Baton, D.L.; de Sauvage, F.J.
A:Title	Genomic structure, chromosomal localization, and conserved alternative splice f
A:Reference	number: I52610; MUID:95152076; PMID:7849319
A:Accession	I52610
A:Status	preliminary; translated from GB/EMBL/DBJ
A:Molecule type	DNA
A:Residues	1-353 <RE>
RESULT 14	
AE0959	solute binding receptor protein [imported] - <i>Salmonella enterica</i> subsp. <i>enterica</i> serovar Typhi
A:Species	<i>Salmonella enterica</i> subsp. <i>enterica</i> serovar Typhi
A:Note	this species has also been called <i>Salmonella</i> typhi
C:Date	09-Nov-2001 #sequence_revision 09-Nov-2001 #text_change 18-Nov-2002
C:Accession	AE0959
R:Parkhill	J.; Dougan, G.; James, K.D.; Thomson, N.R.; Pickard, D.; Wain, J.; Churcher, S.; Maiden, M.C.; Holden, M.T.G.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, K.; Whitehead, S.; Barrell, B.; Parkhill, J.; Wren, B.W.; Thomson, N.R.; Titball, R.W.; Holden, M.T.G.; Prentice, M.B.; demo-Tarraga, A.M.; Chillingworth, T.; Cronin, A.; Davies, R.M.; Davis, P.; Dougan, G.; IL, M.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, K.; Whitehead, S.; Barrell, B.; Nature 413, 523-527, 2001
A:Title	Genome sequence of <i>Yersinia pestis</i> , the causative agent of plague.
A:Reference	number: AB0001; MUID:21470413; PMID:11586360
A:Accession	AB0001
A:Status	preliminary
A:Molecule type	DNA
A:Residues	1-323 <KDR>
A:Cross-references	UNIPROT:Q8ZDC8; UNIPARC:UPI0000DCAA2; GB:AL590842; PIDN: CAC2889.1
C:Genetics	
C:Gene	nrdF
C:Superfamily	ribonucleoside-diphosphate reductase
C:Keywords	oxidoreductase
Query Match	Best Local Similarity 25.2%; Pred. No. 0.94; Score 88; DB 2; Length 323; Matches 34; Conservative 20; Mismatches 59; Indels 22; Gaps 5;
QY	38 NTWTPDKTUNFKYAWKRMVGQDAVEVWVQGLALLSEAVLRRGQALLVNSQWPWPLQVND- 96
Db	2 NTVKPRITPSAISWNKIE-DDDKDEVVN--RITSNFWPEKPLSNPDSNIPSWMTLPBHQ 58
QY	97 ---KAVSGSLSLTLLRAIGAO---KEAISPPDAASAPLRTTADTRKLUFVSYLERGKLK 150
Db	59 LTTRVFTGTTLQIOTNGQAPALIKOAITPHEEAISNISMEAVARSVISIPL 116
QY	151 GKRLKTYGCACTGD 165
Db	117 -----CLTS 121

, S.; Moule, S.; O'Gaora, P.
 Nature 413, 848-852, 2001
 A; Authors: Parry, C.; Quail, M.; Rutherford, K.; Simmonds, M.; Skelton, J.; Stevens, K.;
 A; Reference number: AB0502; MUID:21534947; PMID:11677608
 A; Accession: AE0959
 A; Status: preliminary
 A; Molecule type: DNA
 A; Residues: 1-346 <PAR>
 A; Cross-references: UNIPARC:UPI00005A6A5; GB:AL513382; PIDN:CAD03169.1; PID:916504804;
 C; Genetics:
 A; Gene: STY3352

Query Match 10.3%; Score 87.5; DB 2; Length 346;
 Best Local Similarity 26.7%; Pred. No. 1.1;
 Matches 44; Conservative 22; Mismatches 48; Indels 51; Gaps 9;
 QY 10 RVLERPVLLKEAENITG-CABHCSLMR--NITVPTDTKVNFKWAKRMEVGQAAVEWQ 65
 Db 217 RNLUQEMLEKHPDANVAGSIAAEEAMGRNLITPLTIVSPFL-----THQYR 267
 QY 66 GLALISeAVLRGQALIVNSP-PWEPLOLHQDVKAVSGLISLTILRAQKA-EKAISP 122
 Db 268 GLK-----RGHTILMASSQMANW-----GELAITOSISKVQGQPVDENIIRR 309
 QY 123 -----DAASAPLRTITADTFRKLPFRVVSNFRLGKLUYTGEA 160
 Db 310 VLILTHNNADSVARRSLSPPGFRPVY-----LYQYTSIA 344

RESULT 15

megakaryocyte growth and development factor, long form - human

N; Alternative names: MPL ligand, long form

C; Species: Homo sapiens (man)
 C; Date: 20-Feb-1995 #sequence_revision 20-Feb-1995 #text_change 07-May-1999

C; Accession: A55330
 R; Chang, M.; McNinch, J.; Basu, R.; Shutter, J.; Hsu, R.; Perkins, C.; Mar, V.; Suggs, S.

J. Biol. Chem. 270, 511-514, 1995
 A; Title: Cloning and characterization of the human megakaryocyte growth and development

A; Reference number: A55330; MUID:95122483; PMID:7822271

A; Accession: A55330
 A; Status: preliminary; not compared with conceptual translation

A; Molecule type: DNA
 A; Residues: 1-286 <CHA>

A; Cross-references: UNIPARC:UPI0000148CB2; GB:U17071

C; Genetics:
 A; Gene: MGDP
 A; Map position: 3q26.3

C; Keywords: alternative splicing; cytokine

Query Match 10.2%; Score 86; DB 2; Length 286;
 Best Local Similarity 26.6%; Pred. No. 1.3;
 Matches 41; Conservative 18; Mismatches 75; Indels 20; Gaps 5;

Db 1 APPRLICDSRVLERYLTLAEKAENITGCKEHCSSLNTITVPTDTKVNFKWAKRMEVGQAA 60
 Db 24 APP--ACDURVLSKLIRDHSRLHSRLSQCPVEHPLPPTPVLPVADPFSLGCEWKTQMEETKA 81
 QY 61 VEVMQGLALISeAVLRGQALIVNSP-PWEPLOLHQDVKAVSGLISLTILRAQKA-EKAISP 118
 Db 82 QDILGAVTLLLEGVMARGQGQGPTCLSSLGQLGQSGQVRLLGALQSL-----LGTQ--- 132
 QY 119 ISPPDRAASAPLRTTADTPRKLFRVYSNPLRGK 152
 Db 133 -LPPQG-----RITPAHKDDNAIFLSFQHHLRGK 159

Search completed: February 28, 2006, 15:28:24
 Job time : 41 Secs

GenCore version 5.1.7
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OM protein - protein search, using sw model

Run on: February 28, 2006, 15:20:40 ; Search time 229 Seconds
508.350 Million cell updates/sec

Title: US-10-706-701-1
Perfect score: 846
Sequence: 1 APPRLICDSRVLLERYLREAK. SNFLURGKLUKYGRACRNGD 165

Scoring table: BLOSUM62
Gapop 10.0 , Gapext 0.5

Searched: 2166443 seqs, 705528306 residues

Total number of hits satisfying chosen parameters: 2166443

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

Database : UniProt_05.80;*
1: uniprot_sprot:
2: uniprot_trembl:
Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query	Match Length	DB ID	Description
1	846	100.0	193	1	BPO_HUMAN
2	846	100.0	193	2	Q549U2_HUMAN
3	764.5	90.4	192	1	EPO_MACMO
4	759.5	89.8	192	1	EPO_HORSE
5	723	85.5	192	1	EPO_FELICA
6	705	83.5	192	1	EPO_RAT
7	701	82.9	192	1	BPO_CANFA
8	693	81.9	206	1	BPO_BOVINA
9	692.5	81.4	192	1	BPO_MOUSE
10	689	81.0	194	1	BPO_SHEEP
11	685.5	81.0	195	1	BPO_RABIT
12	680	80.4	192	2	Q6HSS9_GRODE
13	678	80.1	192	2	Q6HBT0_SPADW
14	678	80.1	192	2	Q6HBT1_GRODE
15	678	80.1	192	1	BPO_PIG
16	678	80.1	194	1	B6HBT2_GRODE
17	674	79.7	192	2	Q6HBT3_GRODE
18	663	78.4	133	2	Q8HZ88_SPDRM
19	658	77.8	133	2	Q8HZ89_PANTR
20	627	74.1	131	2	Q8HZ87_PONPY
21	607	71.7	133	2	Q8HZ86_QPRMY
22	554	65.5	133	2	Q8HZ85_SAGOB
23	250	29.6	185	2	Q5LGQ0_EPICG
24	241	28.5	180	2	Q4T554_TETNG
25	241	28.5	195	2	Q6UM11_TETNG
26	238	28.1	182	2	Q6JY23_FOGRU
27	238	28.1	2	Q6YV22_FUGRU	
28	188	22.2	50	2	Q9QV40_9MUR1
29	113	13.4	177	2	Q61VE9_CHICK
30	352	12.9	1	TPO_CANFA	
31	89	10.5	353	1	TPO_HUMAN

ALIGNMENTS

RESULT 1

ID	BPO_HUMAN	STANDARD;	PRT;	193 AA.
AC	P01583; Q9UDZ0; Q9UZ25; Q9UHA0;			
DT	21-JUN-1986 (Rel. 01, Last sequence update)			
DR	10-MAY-2005 (Rel. 47, Last annotation update)			
DB	Erythropoletin precursor (Epoetin).			
GN	Name=BPO;			
OS	Homo sapiens (Human);			
OC	Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Mammalia; Buteraria; Buarchontoglires; Primates; Catarrhini; Hominoidea; Homo.			
OX	NCBI_TAXID=9606;			
RN	[1] MEDLINE=85137839; PubMed=3838366;			
RX	NUCLEOTIDE SEQUENCE.			
RA	Jacobs K., Shoemaker C., Rudersdorf R., Neill S.D., Kaufman R.J., Mufson A., Seehra J., Jones S.S., Hewick R., Fritsch E.F., Kawakita M., Shimizu T., Miyake T., Shima T., Martin F., Stabinsky Z., Badrawi S.M., Lai P.-H., Chen K.K., Fox G.M., Goldwasser E.; "Isolation and characterization of genomic and cDNA clones of human erythropoietin;" Nature 313:806-810(1985).			
RL	[2] MEDLINE=86067948; PubMed=3865178;			
RN	RX NUCLEOTIDE SEQUENCE.			
RA	Lin F.-K., Suggs S., Lin C.-H., Browne J.K., Smalling R., Earie J.C., Chen K.K., Fox G.M., Martin F., Stabinsky Z., Badrawi S.M., Lai P.-H., Goldwasser E.; "Cloning and expression of the human erythropoietin gene."; Proc. Natl. Acad. Sci. U.S.A. 82:7580-7584(1985).			
RN	[3] MEDLINE=99018118; PubMed=3799793;			
RP	NUCLEOTIDE SEQUENCE.			
RA	Gloeckner G., Scherer S., Schattrevoy R., Boright A.P., Weber J., Tsui L.-C., Rosenthal A.; "Large-scale sequencing of two regions in human chromosome 7q22: analysis of 650 kb of genomic sequence around the EPO and CNTN1 loci reveals 17 genes;" Genome Res. 8:1060-1073(1998).			
RL	[4] MEDLINE=93384593; PubMed=6396923;			
RN	NUCLEOTIDE SEQUENCE OF 58-193, AND VARIANTS HEPATOCELLULAR CARCINOMA			
RA	Rupert J.L., Hochachka P.W.; "Erythropoletin gene sequence in the Quechua, a high altitude native population;" Submitted (NOV-1999) to the EMBL/GenBank/DBJ databases.			
RL	[5] MEDLINE=93384593; PubMed=6396923;			
RN	NUCLEOTIDE SEQUENCE OF 58-193, AND VARIANTS HEPATOCELLULAR CARCINOMA			
RA	Funakoshi A., Muta H., Baba T., Shimizu S.; "Gene expression of mutant erythropoietin in hepatocellular carcinoma;" Biochem. Biophys. Res. Commun. 195:717-722(1993).			
RL				

RN	[6]	PROTEIN SEQUENCE OF 28-193, AND DISULFIDE BONDS.	DR	EMBL; M11319; AAC52400.1; -; Genomic_DNA.
RP		PROTEIN TISSUE-Urine;	DR	EMBL; AF053356; AAC78191.1; -; Genomic_DNA.
RC		RECOMBINANT TISSUE-Urine;	DR	EMBL; AF202308; AAC23132.1; -; Genomic_DNA.
RX		MEDLINE-86140080; PubMed=3949763;	DR	EMBL; AF202306; AAC23132.1; JOINED; Genomic_DNA.
RA		Lai P.H., Everett R., Wang F.F., Arakawa T., Goldwasser B.; "Structural characterization of human erythropoietin."; J. Biol. Chem. 261:3116-3121(1986).	DR	EMBL; AF202307; AAC23132.1; JOINED; Genomic_DNA.
RT		"Isolation of human erythropoietin with monoclonal antibodies."; J. Biol. Chem. 259:2707-2710(1984).	DR	EMBL; AF202310; AAC23133.1; JOINED; Genomic_DNA.
RL		[7]	DR	EMBL; AF202309; AAC23133.1; JOINED; Genomic_DNA.
RN		PRELIMINARY PROTEIN SEQUENCE OF 28-57.	DR	EMBL; AF202311; AAC17572.1; -; Genomic_DNA.
RP		RX	DR	EMBL; AF202314; AAC23134.1; -; Genomic_DNA.
RX		MEDLINE-84135751; PubMed=6689899;	DR	EMBL; AF22312; AAC23134.1; JOINED; Genomic_DNA.
RA		Yanagawa S., Hirade K., Ohnishi H., Sabaki R., Chiba H., Ueda M., Goto M.; "Isolation of human erythropoietin with monoclonal antibodies."; J. Biol. Chem. 259:2707-2710(1984).	DR	EMBL; AF202313; AAC23134.1; JOINED; Genomic_DNA.
RA		[8]	DR	EMBL; S65458; ADJ13964.1; -; mRNA.
RP		STRUCTURE OF CARBOHYDRATES.	DR	PIR; A01855; ZNUO.
RX		MEDLINE-8815657; PubMed=3336214;	DR	PDB; 1BQY; NMR; A=28-193.
RA		Kobata A.; "Comparative study of the asparagine-linked sugar chains of human erythropoietins purified from urine and the culture medium of recombinant Chinese hamster ovary cells."; J. Biol. Chem. 263:3657-3663(1988).	DR	PDB; 1C94; X-ray; C=28-193.
RN		[9]	DR	Glycosuitedb; P01588; -.
RP		STRUCTURE OF CARBOHYDRATES.	DR	Ensembl; ENSG0000130427; Homo sapiens.
RX		MEDLINE-89118279; PubMed=3219367;	DR	HGNC; HGNC:3415; EPO.
RA		Sasaki H., Ochi N., Dell A., Fukuda M.; "Site-specific glycosylation of human recombinant erythropoietin: analysis of glycopeptides or peptides at each glycosylation site by fast atom bombardment mass spectrometry."; Biochemistry 27:8618-8626(1988).	DR	MIM; 131170; -.
RT		[10]	DR	GO; GO:0005615; C:extracellular space; TAS.
RT		RT	DR	GO; GO:0008015; P:circulation; NAS.
RT		RT	DR	GO; GO:0006950; P:response to stress; TAS.
RT		RT	DR	GO; GO:007165; P:signal transduction; NAS.
RL		DR	DR	InterPro; IPRO1231; Cytokine_4_hlx.
RN		DR	DR	InterPro; IPRO01123; EPO_TPO.
RP		STRUCTURE OF CARBOHYDRATES.	DR	InterPro; IPRO03013; Erythropoetin.
RA		MEDLINE-9231463; PubMed=1820196;	DR	PANTHER; PTBL10370; Erythropoetin; 1.
RT		Takeuchi M., Kobata A.; "Structures and functional roles of the sugar chains of human erythropoietins."; Glycobiology 1:337-346(1991).	DR	Pran; PR00758; EPO_TPO; 1.
RL		[11]	DR	PINSF; PIRSF001951; EPO; 1.
RP		X-RAY CRYSTALLOGRAPHY (1.9 ANGSTROMS).	DR	PRINTS; PR00272; ERYTHROPTN.
RX		MEDLINE-98454929; PubMed=9774108; DOI:10.1038/26773;	DR	PROSITE; PS00817; EPO_TPO; 1.
RA		SYED R.S., Reid S.W., Li C., Cheetham J.C., Aoki K.H., Liu B., Zhan H., Ossblund T.D., Chirino A.J., Zhang J., Finer-Moore J., Elliott S., Sitney K., Katz B.A., Matthews D.J., Wendoloski J.J., RA	DR	KW
RA		Egrie J., Stroud R.M., "Efficiency of signalling through cytokine receptors depends critically on receptor orientation."; Nature 395:511-516(1998).	FT	Glycoprotein; Hormone; Pharmaceutical; Polymorphism; Signal.
RT		-1- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.	FT	CHAIN
CC		-1- SUBCELLULAR LOCATION: Secreted.	FT	28 193
CC		-1- TISSUE SPECIFICITY: Produced by kidney or liver of adult mammals and by liver of fetal or neonatal mammals.	FT	PROPER
CC		-1- PHARMACEUTICAL: Used for the treatment of anemia. Available under the name EpoGen (Amgen), EpoGen (Chugai), EpoMax (Elanex), Eprex (Janssen-Cilag), Neorecombin (Roche), and Procrit (Ortho Biotech). Variations in the glycosylation pattern of EPO distinguishes these products. EpoGen, EpoGen, Eprex and Procrit are generically known as epoetin alfa, Neorecombin and Recombin AB epoetin beta and EpoMax as epoetin omega.	FT	CARBOHYD
CC		-1- SIMILARITY: Belongs to the EPO/TFP family.	FT	51 51
CC		-1- DATABASE: NAME=R&D Systems' cytokine source book: EPO; WWW=http://www.rndsystems.com/aspx/gsitebuilder.asp?bodyId=197".	FT	CARBOHYD
CC		This Swiss-Prot entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the EMBL outstation - the European Bioinformatics Institute. There are no restrictions on its use as long as its content is in no way modified and this statement is not removed.	FT	DISULFID
CC		DR EMBL; X02158; CAA2095.1; -; Genomic_DNA.	FT	VARIANT
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	149 149
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	CONFLICT
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	85 85
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	COFLICT
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	140 140
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	HELIK
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	32 34
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	52 55
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	53 55
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	57 58
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	61 68
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	73 73
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	75 78
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	79 80
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	83 109
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	118 138
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	139 140
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	141 147
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	148 149
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	160 164
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	165 177
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	178 178
CC		EMBL; X02157; CAA2094.1; -; mRNA.	FT	179 188
DR		EMBL; X02157; CAA2094.1; -; mRNA.	SQ	SEQUENCE
DR		EMBL; X02157; CAA2094.1; -; mRNA.	SQ	193 AA; 21307 MM; C91FB54C26A52033 CRC64;

Query Match 100.0%; Score 846; DB 1; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.2e-72;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKAENTTGCABHSLENNTVPTDKTPNYKAWRMVEGQA 60
 DR 28 APPRLICDSRVLERYLLEAKAENTTGCABHSLENNTVPTDKTPNYKAWRMVEGQA 87
 RC NIH MGC Project;
 RG Submitted (APR-2005) to the EMBL/GenBank/DDJB databases.
 RL CC -1- SUBCELLULAR LOCATION: Secreted (BY Similarity).
 RP NUCLEOTIDE SEQUENCE.
 TISSUE=Brain;
 DR EMBL; AC009488; AAP22357.1; -; Genomic_DNA.
 QY 61 VEWQGLALLSEAVLRGQALVNSSOPWEPLQHDKAVSGLSLTLLRAIGAQEAI 120
 DR SWR; Q54902; 28-193.
 DR 88 VEWQGLALLSEAVLRGQALVNSSOPWEPLQHDKAVSGLSLTLLRAIGAQEAI 147
 DR GO; GO:0005576; C:extracellular region; IEA.
 DR GO; GO:0005128; F:erythropoietin receptor binding; IEA.
 DR GO; GO:0005179; F:hormone activity; IEA.
 DR IPR012351; Cyrotope_4_hlx.
 DR IPR01323; EPO_TPO.
 DR InterPro; IPR003013; Erythropoietin.
 DR Pfam; PF00758; EPO_TPO; 1.
 DR PRINTS; PRO227; ERYTHROPTN.
 DR PROSTK; PS0087; EPO_TPO; 1.
 DR Hormone; Hypothetical_protein.
 RN 13-SEP-2005 (TREMBL); 31; Last sequence update)
 DT 13-SEP-2005 (TREMBL); 31; Last annotation update)
 DR Hypothetical protein EPO (Erythropoietin,).
 DR Name=EGP0;
 DR Homo sapiens (Human);
 OC Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Butheria; Euarchontoglires; Primates; Catarrhini; Hominoidea;
 OC Homo; Homo sapiens (Human);
 OX NCBI_TAXID=9606;
 RN [1]
 RP NUCLEOTIDE SEQUENCE.
 RX MEDLINE=90063792; PubMed=9847074;
 RA Wilson R.;
 RT "Toward a complete human genome sequence.";
 RL Genome Res. 8:1097-1108(1998).
 RN [2]
 RP NUCLEOTIDE SEQUENCE.
 RA Doebeke A., Elliott G., Jones T., Nguyen C., Stoneking T., Sun H.;
 RT "The sequence of Homo sapiens BAC clone RP11-336D7.";
 RL Submitted (AUG-1999) to the EMBL/GenBank/DDJB databases.
 RN [3]
 RA Waterston R.H.;
 RT NUCLEOTIDE SEQUENCE.
 RL Submitted (MAY-2001) to the EMBL/GenBank/DDJB databases.
 RN [4]
 RP NUCLEOTIDE SEQUENCE.
 RA Waterston R.;
 RL Submitted (APR-2003) to the EMBL/GenBank/DDJB databases.
 RN [5]
 RP NUCLEOTIDE SEQUENCE.
 RC TISSUE=Brain;
 MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
 RA Strausberg R.L., Feingold B.A., Grouse L.H., Derge J.G.,
 RA Klausner R.D., Collins F.S., Wagner L., Shemesh C.M., Schuler G.D.,
 RA Altschul S.F., Zeeberg B.B., Bustow K.H., Schaefer C.F., Bhat N.K.,
 RA Hopkins R.P., Jordan H., Moore T., Max S.I., Wang J., Heih F.,
 RA Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
 RA Ditchkenko D., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
 RA Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Schetetz T.B.,
 RA Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
 RA Raha S.S., Loquaiello N.A., Peters G.J., Abramson R.D., Mulahay S.J.,
 RA Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
 RA Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
 RA Fahey J., Helton B., Kettman M., Madan A., Rodriguez S., Sanchez A.,
 RA Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
 RA Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
 RA Rodriguez A.C., Grimes J., Schmutz J., Myers R.M.,
 RA Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.B.,
 RA Schenck A., Schein J.E., Jones S.J.M., Marra M.A.;
 RT "Generation and initial analysis of more than 15,000 full-length human
 and mouse cDNA sequences.";
 RT and mouse cDNA sequences.";

RESULT 2
 Q54902_HUMAN
 ID Q54902_HUMAN PRELIMINARY; PRT; 193 AA.
 DT 13-SEP-2005 (TREMBL); 31; Last sequence update)
 DT 13-SEP-2005 (TREMBL); 31; Last annotation update)
 DR Hypothetical protein EPO (Erythropoietin,).
 DR Name=EGP0;
 DR Homo sapiens (Human);
 OC Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Butheria; Euarchontoglires; Primates; Catarrhini; Hominoidea;
 OC Homo; Homo sapiens (Human);
 OX NCBI_TAXID=9606;
 RN [1]
 RP NUCLEOTIDE SEQUENCE.
 RX MEDLINE=90063792; PubMed=9847074;
 RA Wilson R.;
 RT "Toward a complete human genome sequence.";
 RL Genome Res. 8:1097-1108(1998).
 RN [2]
 RP NUCLEOTIDE SEQUENCE.
 RA Doebeke A., Elliott G., Jones T., Nguyen C., Stoneking T., Sun H.;
 RT "The sequence of Homo sapiens BAC clone RP11-336D7.";
 RL Submitted (AUG-1999) to the EMBL/GenBank/DDJB databases.
 RN [3]
 RA Waterston R.H.;
 RT NUCLEOTIDE SEQUENCE.
 RL Submitted (MAY-2001) to the EMBL/GenBank/DDJB databases.
 RN [4]
 RP NUCLEOTIDE SEQUENCE.
 RA Waterston R.;
 RL Submitted (APR-2003) to the EMBL/GenBank/DDJB databases.
 RN [5]
 RP NUCLEOTIDE SEQUENCE.
 RC TISSUE=Brain;
 MEDLINE=22388257; PubMed=12477932; DOI=10.1073/pnas.242603899;
 RA Strausberg R.L., Feingold B.A., Grouse L.H., Derge J.G.,
 RA Klausner R.D., Collins F.S., Wagner L., Shemesh C.M., Schuler G.D.,
 RA Altschul S.F., Zeeberg B.B., Bustow K.H., Schaefer C.F., Bhat N.K.,
 RA Hopkins R.P., Jordan H., Moore T., Max S.I., Wang J., Heih F.,
 RA Richards S., Worley K.C., Hale S., Garcia A.M., Gay L.J., Hulyk S.W.,
 RA Ditchkenko D., Marusina K., Farmer A.A., Rubin G.M., Hong L.,
 RA Stapleton M., Soares M.B., Bonaldo M.F., Casavant T.L., Schetetz T.B.,
 RA Brownstein M.J., Usdin T.B., Toshiyuki S., Carninci P., Prange C.,
 RA Raha S.S., Loquaiello N.A., Peters G.J., Abramson R.D., Mulahay S.J.,
 RA Bosak S.A., McEwan P.J., McKernan K.J., Malek J.A., Gunaratne P.H.,
 RA Villalon D.K., Muzny D.M., Sodergren E.J., Lu X., Gibbs R.A.,
 RA Fahey J., Helton B., Kettman M., Madan A., Rodriguez S., Sanchez A.,
 RA Whiting M., Madan A., Young A.C., Shevchenko Y., Bouffard G.G.,
 RA Blakesley R.W., Touchman J.W., Green E.D., Dickson M.C.,
 RA Rodriguez A.C., Grimes J., Schmutz J., Myers R.M.,
 RA Butterfield Y.S.N., Krzywinski M.I., Skalska U., Smailus D.B.,
 RA Schenck A., Schein J.E., Jones S.J.M., Marra M.A.;
 RT "Generation and initial analysis of more than 15,000 full-length human
 and mouse cDNA sequences.";

RESULT 3
 EPO_MACPA STANDARD; PRT; 192 AA.
 ID EPO_MACPA
 AC P07865;
 DT 01-AUG-1988 (Rel. 08; Created)
 DT 01-AUG-1989 (Rel. 08; Last sequence update)
 DT 10-MAY-2005 (Rel. 47; Last annotation update)
 DR Erythropoietin precursor.
 DR Name=EGP0;
 DR Macaca fascicularis (Crab eating macaque) (Cynomolgus monkey).
 OC Bukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi;
 OC Mammalia; Butheria; Euarchontoglires; Primates; Catarrhini;
 OC Cercopithecoidea; Cercopithecinae; Macaca.
 OC NCBI_TAXID=9541;
 RN [1]
 RP NUCLEOTIDE SEQUENCE.
 RX MEDLINE=87055236; PubMed=2877922; DOI=10.1016/0378-1119(86)90183-6;
 RA Lin F.-K., Lin C.-H., Lai P.-H., Browne J.K., Egrie J.C., Smalling R.,
 RA Fox G.M., Chen K.K., Castro M., Sugars S.;
 RT "Monkey erythropoietin gene: cloning, expression and comparison with
 the human erythropoietin gene.";
 Gene 4:201-203(1986).
 CC -1- FUNCTION: Erythropoietin is the principal hormone involved in the
 regulation of erythrocyte differentiation and the maintenance of a
 physiological level of circulating erythrocyte mass.
 CC -1- SUBCELLULAR LOCATION: Secreted.
 CC -1- TISSUE SPECIFICITY: Produced by kidney or liver of adult mammals
 and by liver of fetal or neonatal mammals.
 CC -1- SIMILARITY: Belongs to the EPO/TPO family.
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CC EMBL; M18189; AAA36841.1; - ; mRNA.
CC PIR; J00173; J00173.
DR HSSP; P01588; 1C14.
DR SIR; P07865; 28-192.
DR InterPro; IPR012351; Cytokine_4_hlx.
DR InterPro; IPR001323; EPO_TPO.
DR InterPro; IPR03013; Erythropoetin.
DR PANTHER; PTHR10370; Erythropoetin; 1.
DR Pfam; PF00758; EPO_TPO; 1.
DR PIRSF; PIRSF001951; EPO; 1.
DR PRINTS; PRO0072; ERYTHROPTN.
DR PROSITE; PS00817; EPO_TPO; 1.
DR PRINTS; PRO0072; ERYTHROPTN.
KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.
FT SIGNAL 1 27
FT CHAIN 28 192
FT CARBOHYD 51 51
FT CARBOHYD 65 65
FT CARBOHYD 110 110
FT CARBOHYD 152 152
FT DISULFID 34 187
FT DISULFID 56 60
SQ SEQUENCE 192 AA; 21114 MW; BBA900B442AD4522 CRC64;

Query Match 90.4%; Score 764.5; DB 1; Length 192;
Best Local Similarity 91.5%; Pred No. 1-3e-64;
Matches 151; Conservative 7; Mismatches 6; Indels 1; Gaps 1;

Qy 1 APPRLICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNYAWKMEVQQA 60
Db 28 APPRLICDSRVLYLEAKEAENITVPGCSECSLENITVPDKVNYAWKMEVQQA 87

Qy 61 VEWQGLALISEAVLRGQLLVNSQPWPEPLQHVAKGAVSGRSLTTLRGAQEAIS 120
Db 88 VEWQGLALISEAVLRGQLLVNSQPWPEPLQHVAKGAVSGRSLTTLRGAQ-EAIS 146

Db 147 LPDARSAAPLRTTADTFCKLFRVYNSFLRGKLUKYGEACRGD 191

RESULT 4

EPO_MACMU STANDARD; PRT; 192 AA.

ID QP8513; 01-NOV-1997 (Rel. 35, Created)
DT 10-MAY-2005 (Rel. 47, Last annotation update)
DB Erythropoietin precursor.
GN Name=EPO;
OS Macaca mulatta (Rhesus macaque).
OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
OC Mammalia; Eutheria; Euarchontoglires; Primates; Catarrhini;
OC Cercopithecidae; Cercopithecoinae; Macaca.
NCBI_TaxID=9544;
[1] NUCLEOTIDE SEQUENCE.
Tissue=Kidney;
RX MEDLINE=93372347; PubMed=8364201;
RA Wen D., Boissel J.-P.R., Tracy T.E., Gruninger R.H., Mulcahy L.S.,
RA Czaplinski J., Goodman M., Bonn H.F.;
RT "Erythropoietin structure-function relationships: high degree of
RT sequence homology among mammals";
RL Blood 82:1507-1516(1993);
CC -- FUNCTION: Erythropoietin is the principal hormone involved in the
regulation of erythrocyte differentiation and the maintenance of a
physiological level of circulating erythrocyte mass.
CC -- SUBCELLULAR LOCATION: secreted.
CC -- TISSUE SPECIFICITY: Produced by kidney or liver of adult mammals
and by liver of fetal or neonatal mammals.
CC -- SIMILARITY: Belongs to the EPO/TPO family.

CC -----
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CC between the Swiss Institute of Bioinformatics and the EMBL outstation -
the European Bioinformatics Institute. There are no restrictions on its
use as long as its content is in no way modified and this statement is not
removed.
CC -----
CC EMBL; LI0609; AAA36842.1; - ; mRNA.
CC PIR; I04613; I86613.
DR HSSP; P01588; 1C14.
DR SIR; Q28513; 28-192.
DR InterPro; IPR013351; Cytokine_4_hlx.
DR InterPro; IPR001323; EPO_TPO.
DR InterPro; IPR03013; Erythropoetin.
DR PANTHER; PTHR10370; Erythropoetin; 1.
DR Pfam; PF00758; EPO_TPO; 1.
DR PIRSF; PIRSF001951; EPO; 1.
DR PRINTS; PRO0072; ERYTHROPTN.
DR PROSITE; PS00817; EPO_TPO; 1.
DR PRINTS; PRO0072; ERYTHROPTN.
KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.
FT SIGNAL 1 27
FT CHAIN 28 192
FT CARBOHYD 51 51
FT CARBOHYD 65 65
FT CARBOHYD 110 110
FT CARBOHYD 152 152
FT DISULFID 34 187
FT DISULFID 56 60
SQ SEQUENCE 192 AA; 21081 MW; 27550B264628CD1 CRC64;

Query Match 89.8%; Score 759.5; DB 1; Length 192;
Best Local Similarity 90.3%; Pred No. 4e-64;
Matches 149; Conservative 9; Mismatches 6; Indels 1; Gaps 1;

Qy 1 APPRLICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNYAWKMEVQQA 60
Db 28 APPRLICDSRVLYLEAKEAENITVPGCSECSLENITVPDKVNYAWKMEVQQA 87

Qy 61 VEWQGLALISEAVLRGQLLVNSQPWPEPLQHVAKGAVSGRSLTTLRGAQEAIS 120
Db 88 VEWQGLALISEAVLRGQLLVNSQPWPEPLQHVAKGAVSGRSLTTLRGAQ-EAIS 146

Db 147 LPDARSAAPLRTTADTFCKLFRVYNSFLRGKLUKYGEACRGD 191

RESULT 5

EPO_HORSE STANDARD; PRT; 192 AA.

ID EPO_HORSE AC Q867B1; 10-MAY-2005 (Rel. 47, Created)
DT 10-MAY-2005 (Rel. 47, Last annotation update)
DB Erythropoietin precursor.
GN Name=EPO;
OS Equus caballus (Horse).
OC Mammalia; Eutheria; Laurasiatheria; Perissodactyla; Equidae; Equus.
OX NCBI_TaxID=9196;
[1] NUCLEOTIDE SEQUENCE.
Tissue=Kidney;
RX PubMed=4719696;
RA Saito F., Yamashita S., Kugo T., Hasegawa T., Mitsui I.,
RA Kijima-Suda T.;
RT "Nucleotide sequence of equine erythropoietin and characterization of
region-specific antibodies";
RL Am. J. Vet. Res. 65:15-19(2004).
CC -- FUNCTION: Erythropoietin is the principal hormone involved in the
regulation of erythrocyte differentiation and the maintenance of a
physiological level of circulating erythrocyte mass (By
similarity).

RT "Erythropoietin structure-function relationships: high degree of sequence homology among mammals.";

RT sequence homology among mammals.";

RL Blood 82:1507-1516 (1993).

CC -1- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.

CC -1- SUBCELLULAR LOCATION: Secreted.

CC -1- TISSUE SPECIFICITY: Produced by kidney or liver of adult mammals and by liver of fetal or neonatal mammals.

CC -1- SIMILARITY: Belongs to the EPO/TPO family.

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CC EMBL; ABI0030; BAC55239.1; - ; mRNA.

DR RESP; Q667B1; 27-192.

DR InterPro; IPR012351; Cytokine_4_hlx.

DR InterPro; IPR00133; EPO TPO.

DR InterPro; IPR003013; Erythropoetin.

DR PANTHER; PTHR1030; Erythropoetin; 1.

DR Pfam; PRO0758; EPO_TPO; 1.

DR PRSF; PRSF001951; EPO; 1.

DR PRINTS; PR0072; ERYTHROPTN.

DR PROSITE; PS00817; EPO_TPO; 1.

KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.

FT SIGNAL 1 26 By similarity.

FT CHAIN 27 192 Erythropoietin.

FT CARBOHYD 50 50 N-linked (GlcNAc. . .) (Potential).

FT CARBOHYD 64 64 N-linked (GlcNAc. . .) (Potential).

FT CARBOHYD 109 109 N-linked (GlcNAc. . .) (Potential).

FT DISULFID 33 187 BY similarity.

FT DISULFID 55 59 BY similarity.

SQ 192 AA; 20984 MW; B02D09B09C4F CRC64;

Query Match 85.5%; Score 723; DB 1; Length 192;

Best Local Similarity 84.8%; Pred. No. 1.2e-60; Mismatches 10; Indels 0; Gaps 0;

Matches 140; Conservative 10; MisMatches 15; Del 0; Insert 0;

QY 1 APPRLICDSRVLRVYLLEAKRKAENITTCGACGSLENNTIVPDTKVNFTYAWKRMEVGQAA 60

Db 27 APPRLICDSRVLRVYLLEAKRKAENITTCGACGSLENNTIVPDTKVNFTYAWKRMEVGQAA 86

QY 61 VEWVOGLAISSEAVTLRGQLLVNSQWPFLQLAVDKAVSGLSLTILRALGAQEAKTS 120

Db 87 VEWVOGLAISSEAVTLRGQLLANSQSSETLQLAVDKAVSSLSLTSILRALGAQEAKTS 146

RESULT 6

EPO_FELCA ID EPO_FELCA STANDARD; PRT; 192 AA.

AC P33708; 01-FEB-1994 (Rel. 28, Created)

DT 01-OCT-1996 (Rel. 34, Last sequence update)

DT 10-MAY-2005 (Rel. 47, Last annotation update)

DR Erythropoietin precursor.

Name=EPO;

OS Felis silvestris catus (Cat).

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Mammalia; Eutheria; Laurasiatheria; Carnivora; Fissipedia; Felidae; Felinae; Felis.

OX NEBI_TAXID=6685;

[1] NUCLEOTIDE SEQUENCE [mRNA].

RC TISSUE-Kidney.

RA Goodman R.E.; Bell R.G.;

RT "A feline erythropoietin cDNA, cloned by RT/PCR amplification of kidney derived RNA with hybrid (human/mouse) primers"; Submitted (NOV-1993) to the EMBL/GenBank/DBJ databases. [2]

RX MEDLINE-9332347; Pubmed=8364201;

RX RA Wen D., Boiselle J.-P.R., Tracy T.E., Gruninger R.H., Mulcahy L.S., Czelusniak J., Goodman M., Bunn H.F.,

RT "Erythropoietin structure-function relationships: high degree of sequence homology among mammals.";

RT sequence homology among mammals.";

RL Blood 82:1507-1516 (1993).

CC -1- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.

CC -1- SUBCELLULAR LOCATION: Secreted.

CC -1- TISSUE SPECIFICITY: Produced by kidney or liver of adult mammals and by liver of fetal or neonatal mammals.

CC -1- SIMILARITY: Belongs to the EPO/TPO family.

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CC EMBL; U00685; AAA18282.1; - ; mRNA.

DR EMBL; L10606; AAK30807.1; - ; mRNA.

DR PIR; I16083; I46083.

DR HSPB; P01588; 1BY.

DR SMR; P33708; 27-192.

DR InterPro; IPR012351; Cytokine_4_hlx.

DR InterPro; IPR00133; EPO TPO.

DR PANTHER; PTHR1030; Erythropoetin.

DR Pfam; PRO0758; EPO_TPO; 1.

DR PRSF; PRSF001951; EPO; 1.

DR PRINTS; PR0072; ERYTHROPTN.

DR Erythrocyte maturation; Glycoprotein; Hormone; Signal.

FT SIGNAL 1 26 By similarity.

FT CHAIN 27 192 Erythropoietin.

FT CARBOHYD 50 50 N-linked (GlcNAc. . .) (Potential).

FT CARBOHYD 64 64 N-linked (GlcNAc. . .) (Potential).

FT DISULFID 33 187 BY similarity.

FT DISULFID 55 59 BY similarity.

FT CONFLICT 44 44 G -> B (in Ref. 2).

SQ 192 AA; 20914 MW; 61C5EA0F5B937293 CRC64;

Query Match 83.5%; Score 706; DB 1; Length 192;

Best Local Similarity 83.6%; Pred. No. 5e-59; Mismatches 18; Indels 0; Gaps 0;

Matches 138; Conservative 9; MisMatches 18; Del 0; Insert 0;

QY 1 APPRLICDSRVLRVYLLEAKRKAENITTCGACGSLENNTIVPDTKVNFTYAWKRMEVGQAA 60

Db 27 APPRLICDSRVLRVYLLEAKRKAENITTCGACGSLENNTIVPDTKVNFTYAWKRMEVGQAA 86

QY 61 VEWVOGLAISSEAVTLRGQLLVNSQWPFLQLAVDKAVSGLSLTILRALGAQEAKTS 120

Db 87 VEWVOGLAISSEAVTLRGQLLANSQSSETLQLAVDKAVSSLSLTSILRALGAQEAKTS 146

RESULT 7

EPO_RAT ID EPO_RAT STANDARD; PRT; 192 AA.

AC P29676; P70504; 01-APR-1993 (Rel. 25, Created)

DT 01-APR-1993 (Rel. 25, Last sequence update)

DT 10-MAY-2005 (Rel. 47, Last annotation update)

DE Erythropoietin precursor.

Name=Epo;

OS Rattus norvegicus (Rat).

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Sciurognathi; Muroidea; Muridae; Murinae; Rattus.

OX NEBI_TAXID=1016;

[1]

RESULT 9		EPO_BOVIN	
ID	PRT;	ID	PRT;
AC P48617;		AC P48617;	
DT 01-FEB-1996 (Rel. 33, Created)		DT 01-FEB-1996 (Rel. 33, Last sequence update)	
DE 10-MAY-2005 (Rel. 47, Last annotation update)		DE Erythropoietin precursor.	
GN Name=EPO;		GN Name=EPO;	
OS Bos taurus (Bovine).		OS Bos taurus; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Bokaryota; Mammalia; Eutheria; Laurasiatheria; Cetartiodactyla; Ruminantia; Pecora; Bovidae; Bovinae; Bos.	
OC NCBI_TaxID:9913;		OC NCBI_TaxID:9913;	
RN [1]	NUCLEOTIDE SEQUENCE [MRNA].	RN [1]	NUCLEOTIDE SEQUENCE [MRNA].
RP STRAIN=Boran; TISSUE=Kidney;		RP STRAIN=Boran; TISSUE=Kidney;	
RX MEDLINE=96257233; PubMed=86662865; DOI=10.1016/0378-1119(95)00895-0;		RX MEDLINE=96257233; PubMed=86662865; DOI=10.1016/0378-1119(95)00895-0;	
RA Suliman H.B.; Majiwa P.A.O., Feldman B.F., Mertens B., Logan H.B.; Majiwa P.A.O., Feldman B.F., Mertens B., Lili.;		RA Suliman H.B.; Majiwa P.A.O., Feldman B.F., Mertens B., Logan H.B.; Majiwa P.A.O., Feldman B.F., Mertens B., Lili.;	
RT "Cloning of a cDNA encoding bovine erythropoietin and analysis of its transcription in selected tissues.";		RT "Cloning of a cDNA encoding bovine erythropoietin and analysis of its transcription in selected tissues.";	
RL Gene 171:275-280(1996).		RL Gene 171:275-280(1996).	
CC -!- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.		CC -!- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.	
CC -!- SUBCELLULAR LOCATION: Secreted.		CC -!- SUBCELLULAR LOCATION: Produced by kidney or liver of adult mammals	
CC -!- TISSUE SPECIFICITY: Produced by kidney or liver of fetal or neonatal mammals.		CC -!- TISSUE SPECIFICITY: Produced by kidney or liver of fetal or neonatal mammals.	
CC -!- SIMILARITY: Belongs to the EPO/rpo family.		CC -!- SIMILARITY: Belongs to the EPO/rpo family.	
CC	This Swiss-Prot entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the European Bioinformatics Institute. There are no restrictions on its use as long as its content is in no way modified and this statement is not removed.	CC	This Swiss-Prot entry is copyright. It is produced through a collaboration between the Swiss Institute of Bioinformatics and the European Bioinformatics Institute. There are no restrictions on its use as long as its content is in no way modified and this statement is not removed.
DR EMBL: L41354; AAA41268.1; -; mRNA.		DR EMBL: L41354; AAA41268.1; -; mRNA.	
DR RNSP; P01588; 1CM4.		DR RNSP; P01588; 1CM4.	
DR P8617; 26-192; Cytokine_4_hlx.		DR P8617; 26-192; Cytokine_4_hlx.	
DR InterPro; IPR012351; EPO_TPO.		DR InterPro; IPR012351; EPO_TPO.	
DR IPR01323; EPO_TPO.		DR IPR01323; EPO_TPO.	
DR PANTHER; PTM103013; ERYTHROPTN.		DR PANTHER; PTM103013; ERYTHROPTN.	
DR PFAM; PF00758; EPO_TPO_1.		DR PFAM; PF00758; EPO_TPO_1.	
DR PRSF; PRSF001951; EPO_1.		DR PRSF; PRSF001951; EPO_1.	
DR PRINTS; PR00272; ERYTHROPTN.		DR PRINTS; PR00272; ERYTHROPTN.	
DR PROSITE; PS00817; EPO_TPO_1.		DR PROSITE; PS00817; EPO_TPO_1.	
KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.		KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.	
FT SIGNAL_1_25_Potential.		FT SIGNAL_1_25_Potential.	
FT CHAIN_26_192_Erythropoietin.		FT CHAIN_26_192_Erythropoietin.	
FT CARBOHYD_49_49_N-linked (GlcNAc. . .) (Potential).		FT CARBOHYD_49_49_N-linked (GlcNAc. . .) (Potential).	
FT CARBOHYD_63_63_N-linked (GlcNAc. . .) (Potential).		FT CARBOHYD_63_63_N-linked (GlcNAc. . .) (Potential).	
FT CARBOHYD_108_108_BY similarity.		FT CARBOHYD_108_108_BY similarity.	
FT DISULFID_32_187_BY similarity.		FT DISULFID_32_187_BY similarity.	
SQ 54_58_SEQUENCE_192_AA; 21076 MW; DBC419022FB483A CRC64;		SQ 54_58_SEQUENCE_192_AA; 21076 MW; DBC419022FB483A CRC64;	
Query Match 81.9%; Score 692.5; DB 1; Length 192;		Query Match 81.9%; Score 692.5; DB 1; Length 192;	
Best Local Similarity 83.1%; Pred. No. 9.7e-58; Index 1; Gene 1;		Best Local Similarity 83.1%; Pred. No. 9.7e-58; Index 1; Gene 1;	
CC 1 APRLICDSRIVLERYLLEAKENITGCAHCSENITIPDKTKNFKWERMVQQA 60		CC 1 APRLICDSRIVLERYLLEAKENITGCAHCSENITIPDKTKNFKWERMVQQA 60	
CC 26 APRLICDSRIVLERYLLEAKENITGCAHCSENITIPDKTKNFKWERMVQQA 85		CC 26 APRLICDSRIVLERYLLEAKENITGCAHCSENITIPDKTKNFKWERMVQQA 85	
CC 101 LEVWQGLALLSEAIRTGQALLANASQSETPOLAVDKAVSSLSLTSLLRALGAKEAIS 160		CC 101 LEVWQGLALLSEAIRTGQALLANASQSETPOLAVDKAVSSLSLTSLLRALGAKEAIS 160	
Db 121 PPDA-SAAPLTITADTPKFLRVSNFLRGKLUYTGEACRGD 165		Db 121 PPDA-SAAPLTITADTPKFLRVSNFLRGKLUYTGEACRGD 165	
Db 161 LPBEASPAPLRTFDVTLCKFLRVSNFLRGKLUYTGEACRGD 205		Db 161 LPBEASPAPLRTFDVTLCKFLRVSNFLRGKLUYTGEACRGD 205	
RESULT 10		EPO_MOUSE	
ID	PRT;	ID	PRT;
AC P07321;		AC P07321;	
DT 01-APR-1988 (Rel. 07, Created)		DT 01-APR-1988 (Rel. 07, Last sequence update)	
DE 10-MAY-2005 (Rel. 47, Last annotation update)		DE Erythropoietin precursor.	
GN Name=Epo;		GN Name=Epo;	
OS Mus musculus (Mouse).		OS Mus musculus (Mouse).	
OC Mammalia; Buthesia; Euarchontoglires; Glires; Rodentia; Sciurognathi; Muroidea; Murida; Murinae; Mus.		OC Mammalia; Buthesia; Euarchontoglires; Glires; Rodentia; Sciurognathi; Muroidea; Murida; Murinae; Mus.	
OK NCBI_TaxID=10909;		OK NCBI_TaxID=10909;	
RN [1]	NUCLEOTIDE SEQUENCE [DNA].	RN [1]	NUCLEOTIDE SEQUENCE [DNA].
RP MEDLINE=87039105; PubMed=3773894;		RP MEDLINE=87039105; PubMed=3773894;	
RA Shoemaker C.B.; Mitnick L.D.; McDonald J.D., Lin F.-K., Goldwasser E.; RT "Murine erythropoietin gene: cloning, expression, and human gene homology.";		RA Shoemaker C.B.; Mitnick L.D.; McDonald J.D., Lin F.-K., Goldwasser E.; RT "Murine erythropoietin gene: cloning, expression, and human gene homology.";	
RT Mol. Cell. Biol. 6:849-858(1986).		RT Mol. Cell. Biol. 6:849-848(1986).	
RL [2]	NUCLEOTIDE SEQUENCE.	RL [2]	NUCLEOTIDE SEQUENCE.
RN [3]	MEDLINE=87039104; PubMed=3022133;	RN [3]	MEDLINE=87039104; PubMed=3022133;
RP McDonald J.D., Lin F.-K., Goldwasser E.; RT "Cloning, sequencing, and evolutionary analysis of the mouse erythropoietin gene"; RT Mol. Cell. Biol. 6:842-848(1986).		RP McDonald J.D., Lin F.-K., Goldwasser E.; RT "Cloning, sequencing, and evolutionary analysis of the mouse erythropoietin gene"; RT Mol. Cell. Biol. 6:842-848(1986).	
RC SRRN=1291_SV;		RC SRRN=1291_SV;	
RL MEDLINE=21138439; PubMed=11239002; DOI=10.1093/nar/29.6.1352;		RL MEDLINE=21138439; PubMed=11239002; DOI=10.1093/nar/29.6.1352;	
RA Wilson M.D., Riemer C., Martindale D.W., Schnupf P., Boright A.P., RA Cheung T.L., Hardy D.M., Schwartz S.W., Tsui L.-C., RA Miller W., Koop B.F., RA "Comparative analysis of the gene-dense AChR/TPR2 region on human chromosome 7q22 with the orthologous region on mouse chromosome 5.";		RA Wilson M.D., Riemer C., Martindale D.W., Schnupf P., Boright A.P., RA Cheung T.L., Hardy D.M., Schwartz S.W., Tsui L.-C., RA Miller W., Koop B.F., RA "Comparative analysis of the gene-dense AChR/TPR2 region on human chromosome 7q22 with the orthologous region on mouse chromosome 5.";	
RR Cretien S., Duprez V., Maouche L., Gisselbrecht S., Mayeux P., RR Nucleic Acids Res. 29:1352-1365(2001).		RR Cretien S., Duprez V., Maouche L., Gisselbrecht S., Mayeux P., RR Nucleic Acids Res. 29:1352-1365(2001).	
RP NUCLEOTIDE SEQUENCE OF 1-52.		RP NUCLEOTIDE SEQUENCE OF 1-52.	
RC SRRN=ICFW;		RC SRRN=ICFW;	
RL MEDLINE=98030528; PubMed=9365346; DOI=10.1038/sj.onc.1201364;		RL MEDLINE=98030528; PubMed=9365346; DOI=10.1038/sj.onc.1201364;	
RA Chretien S., Duprez V., Maouche L., Gisselbrecht S., Mayeux P., RA "Abnormal erythropoietin (Epo) gene expression in the murine erythroleukemia IW32 cells results from a rearrangement between the G-RT protein beta2 subunit gene and the Epo gene.";		RA Chretien S., Duprez V., Maouche L., Gisselbrecht S., Mayeux P., RA "Abnormal erythropoietin (Epo) gene expression in the murine erythroleukemia IW32 cells results from a rearrangement between the G-RT protein beta2 subunit gene and the Epo gene.";	
RL Oncogene 15:1995-1999(1997).		RL Oncogene 15:1995-1999(1997).	
CC -!- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.		CC -!- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass.	
CC -!- SUBCELLULAR LOCATION: Secreted.		CC -!- SUBCELLULAR LOCATION: Secreted.	
CC -!- TISSUE SPECIFICITY: Produced by kidney or liver of fetal or neonatal mammals.		CC -!- TISSUE SPECIFICITY: Produced by kidney or liver of fetal or neonatal mammals.	
CC -!- SIMILARITY: Belongs to the EPO/TPO family.		CC -!- SIMILARITY: Belongs to the EPO/TPO family.	

OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
 OC Mammalia; Eutheria; Buarchoontoglires; Glires; Lagomorpha; Leporidae;
 OC Oryctolagus;
 OC NCBI_TAXID=9986;

RN [1] NUCLEOTIDE SEQUENCE [GENOMIC DNA / mRNA].
 RC STRAIN>New Zealand white; TISSUE=Kidney.
 RX MEDLINE=21290682; PubMed=11396976; DOI=10.1006/bbrc.2001.5028;
 RA Vilalta A.; Wu D.; Margalit M.; Hobart P.;
 RT "Rabbit EPO gene and cDNA: expression of rabbit EPO after
 intramuscular injection of pDNA."
 RL Biochem Biophys Res Commun. 284:823-827(2001).
 CC -I- FUNCTION: Erythropoietin is the principal hormone involved in the
 regulation of erythrocyte differentiation and the maintenance of a
 physiological level of circulating erythrocyte mass (By
 similarity).
 CC -I- SUBCELLULAR LOCATION: Secreted.
 CC -I- SIMILARITY: Belongs to the EPO/TPO family.

CC This Swiss Prot entry is copyright. It is produced through a collaboration
 CC between the Swiss Institute of Bioinformatics and the EMBL outstation -
 CC the European Bioinformatics Institute. There are no restrictions on its
 CC use as long as its content is in no way modified and this statement is not
 CC removed.

CC

DR EMBL; AF290543; ARG36961.1; ; mRNA.
 DR PIR; JCT699; JCT699.
 DR HSSP; P01588; ICM4.
 DR SNR; OGK2; 29-195.
 DR InterPro; IPR01251; Cytokine_4_hlx.
 DR InterPro; IPR001323; EPO TPO.
 DR InterPro; IPR003013; Erythropoetin.
 DR PANTHER; PTHR10370; Erythropoetin; 1.
 DR Pfam; PF00758; EPO TPO; 1.
 DR DIRSP; PIRSP001951; EPO; 1.
 DR PRINTS; PRO0272; ERYTHROPTN.
 DR PROSITE; PS00817; EPO TPO; 1.
 KW Erythrocyte maturation; Glycoprotein; Hormone; Signal.
 FT SIGNAL 1 28 Potential.
 FT CHAIN 29 195 Erythropoietin.
 FT CARBOHYD 52 52 N-linked (GlcNAc. . .) (Potential).
 FT CARBOHYD 66 66 N-linked (GlcNAc. . .) (Potential).
 FT CARBOHYD 111 111 N-linked (GlcNAc. . .) (Potential).
 FT DISULFID 35 190 By similarity.
 FT DISULFID 57 61 V->A (in ref. 1; AAC36962).
 FT CONFLICT 3 3 SEQUENCE 195 AA: 21054 MW: 09990d7D852713P3 CRC64;

Query Match 80.4%; Score 680.5; DB 1; Length 195;
 Best Local Similarity 81.3%; Pred. No. 1; 4e-56;
 Matches 135; Conservative 12; Mismatches 18; Indels 1; Gaps 1;

QY 1 APPRLICDSRVRVYLRLRAKEAENITTCGAECISLENNTIVPTDKVNFKWAKMEVSGQA 60
 29 APPRLICDSRVRVYLRLRAKEAENITTCGAECISLENNTIVPTDKVNFKWAKMEVSGQA 88

QY 61 VEWMOGLALLSEAVRLQALVLNSSQPWPLOLHVDKAVGSLTSLLRALGAQKAEIS 120
 89 VEWMOGLALLSEAVRLQALVLNSSQPWPLOLHVDKAVGSLTSLLRALGAQKAEIS 148

QY 121 PPDA-SAPLRITTADEPRKLFRVYSNPLRGKLUYGEACRGD 165
 149 PPEASSAALPRTVAADTCKLFRISNPLRGKLUYGEACRGD 194

RESULT 13
 06H89_9RODE 06H89_9RODE PRELIMINARY; PRT; 192 AA.

Db QH8TO SP8JD QH8TO SP8JD PRELIMINARY; PRT; 192 AA.

AC OGH8TO_SP8JD PRELIMINARY; PRT; 192 AA.

AC OGH8TO_SP8JD PRELIMINARY; PRT; 192 AA.

DT 05-JUL-2004 (TREMBLrel. 27, Created)
 DT 05-JUL-2004 (TREMBLrel. 27, Last sequence update)
 DT 05-JUL-2004 (TREMBLrel. 27, Last annotation update)

DE Erythropoietin precursor.
 GN Name=epo;
 OS Spalax galili.
 OC Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
 OC Mammalia; Eutheria; Buarchoontoglires; Glires; Rodentia; Sciurognathi;
 OC Muridae; Spalacinae; Spalax.
 OC NCBI_TAXID=164323;

RN [1] NUCLEOTIDE SEQUENCE.
 RA Shams I.; Avivi A.; Nevo E.;
 RT "Hypoxic stress tolerance of the subterranean mole rat: Expression of
 erythropoietin and hypoxia-inducible factor-1α.";
 RL Nucleic Acids Res. 0:0-0(2004).
 RN [2] NUCLEOTIDE SEQUENCE.
 RC TISSUE=Liver;
 RX TISSUE=Liver;
 RA Shams I.; Avivi A.; Bvaiat N.;
 RT "Hypoxic stress tolerance of the blind subterranean mole rat:
 expression of erythropoietin and hypoxia-inducible factor 1 alpha.";
 RL Proc. Natl. Acad. Sci. U.S.A. 101:9698-9703(2004).
 CC -I- FUNCTION: Erythropoietin is the principal hormone involved in the
 regulation of erythrocyte differentiation and the maintenance of a
 physiological level of circulating erythrocyte mass (By
 similarity).
 CC -I- SUBCELLULAR LOCATION: Secreted (By similarity).
 DR EMBL; AJ71579; CAG2400.1; ; Genomic_DNA.
 DR SNR; Q6H8S9; 27-192.
 DR GO; GO:0005128; F:erythropoietin receptor binding; IEA.
 DR GO; GO:0005179; F:hormone activity; IEA.
 DR InterPro; IPR001323; EPO TPO.
 DR InterPro; IPR003013; Erythropoetin.
 DR PANTHER; PTHR10370; Erythropoetin; 1.
 DR Pfam; PF00758; EPO TPO; 1.
 DR PIRSF; PIRSP001951; EPO; 1.
 DR PRINTS; PRO0272; ERYTHROPTN.
 DR PROSITE; PS00817; EPO TPO; 1.
 KW Erythrocyte maturation; Hormone; Signal.
 FT SIGNAL 1 7 Potential.
 FT CHAIN 8 192 Erythropoietin.
 SO SEQUENCE 192 AA: 21372 MW: 72FC194DBBC5AAB5 CRC64;

Query Match 80.1%; Score 678; DB 2; Length 192;
 Best Local Similarity 80.6%; Pred. No. 2; 3e-56;
 Matches 133; Conservative 8; Mismatches 24; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRVYLRLRAKEAENITTCGAECISLENNTIVPTDKVNFKWAKMEVSGQA 60
 27 APPRLICDSRVRVYLRLRAKEAENITTCGAECISLENNTIVPTDKVNFKWAKMEVSGQA 86

QY 61 VEWMOGLALLSEAVRLQALVLNSSQPWPLOLHVDKAVGSLTSLLRALGAQKAEIS 120
 87 VEWMOGLALLSEAVRLQALVLNSSQPWPLOLHVDKAVGSLTSLLRALGAQKAEIS 146

QY 121 PPDA-SAPLRITTADEPRKLFRVYSNPLRGKLUYGEACRGD 165
 147 PPDTQVPLRRTVDTCFLFRISNPLRGKLUYGEACRGD 191

OC Muridae; Spalacinae; Spalax.
 RA OC
 RT NUBI-TaxID=134510;
 RT NUCLEOTIDE SEQUENCE.
 RC TISSUE=Liver;
 RA Shams I., Avivi A., Nevo E.;
 RA "Hypoxic stress tolerance of the subterranean mole rat: Expression of erythropoietin and hypoxia-inducible factor-1α.";
 RA Nucleic Acids Res. 0:0-0(2004).
 RT NUCLEOTIDE SEQUENCE.
 RC TISSUE=Liver;
 RA Shams I., Avivi A., Nevo E.;
 RA "Hypoxic stress tolerance of the blind subterranean mole rat: expression of erythropoietin and hypoxia-inducible factor 1 alpha.";
 RA Proc. Natl. Acad. Sci. U.S.A. 101:9638-9703(2004).
 -i- FUNCTION: Erythropoietin is the principal hormone involved in the regulation of erythrocyte differentiation and the maintenance of a physiological level of circulating erythrocyte mass (By similarity).
 -i- SUBCELLULAR LOCATION: Secreted (By similarity).
 RT EMBL; AU715794; CAG29399.1; -; Genomic_DNA.
 DR GO; GO:0005576; C:extracellular region; IEA.
 GO; GO:0005128; F:erythropoietin receptor binding; IEA.
 GO; GO:0005179; F:hormone activity; IEA.
 DR InterPro; IPR01323; EPO_TPO.
 DR InterPro; IPR03013; Erythropoetin.
 DR PANTHER; PTHR10370; Erythropoetin; 1.
 DR Pfam; PF00758; EPO_TPO; 1.
 DR PIRSF; PIRSF001951; EPO; 1.
 DR PRINTS; PRO0272; ERYTHROPTN.
 DR KW Erythrocyte maturation; Hormone; Signal.
 FT SIGNAL_1 Potential.
 CHAIN 1 192 AA; 21372 MW; 72FCA94DEBC5AABS CRC64;
 SQ SEQUENCE 192 AA: 21372 MW; 72FCA94DEBC5AABS CRC64;
 Query Match 80.1%; Score 678; DB 2; Length 192;
 Best Local Similarity 80.6%; Pred. No. 2,3e-56;
 Matches 133; Conservative 8; Mismatches 24; Indels 0; Gaps 0;
 QY 1 APPRLICDSVRVLYLKEAENITGCAEHCSLENNTVPDKVNKFYAWKRMVEGQOA 60
 Db 27 APPRLICDSVRVLYLKEAENITGCAEGPFRNFNTVPDKVNKFYAWKTMGVERQA 86
 QY 61 VEWQGLALLSEAVIRGQALLVNSQWPMLQHDKAVSGLSLTTLRAGQKEAIS 120
 Db 87 VEWQGLALLSEAVIRGQALLVNSQWPMLQHDKAVSGLSLTTLRAGQKEAIS 146
 QY 121 PDDASAAPLRTTADTPRKLYVSNFLRGKLYTGACRGED 165
 Db 147 PDDTQVPLRRTVDTCFKLFRYTSNLRGKLYTGACRGED 191
 QY 147 PDDTQVPLRRTVDTCFKLFRYTSNLRGKLYTGACRGED 191
 DB
 GN Name=epo;
 OS Spalax carmeli.
 OC Bathyergidae; Metazoa; Chordata; Craniata; Vertebrata; Buteleostomi;
 OC Mammalia; Buteraria; Euarchontoglires; Glires; Rodentia; Sciurognathi;
 OC Muridae; Spalacinae; Spalax.
 OC NCBI_TaxID=164324;
 RN [1]
 NUCLEOTIDE SEQUENCE.
 TISSUE=Liver;

GenCore version 5.1.7
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OM protein - protein search, using sw model
 Run on: March 1, 2006, 10:19:31 ; Search time 187 Seconds
 (without alignments)
 387.687 Million cell updates/sec

Title: US-10-706-701-1
 Perfect score: 846

Sequence: 1 APPRLICDSRVLERYLVEAKR... SNFLRKGKLUKYTGACRQD 165

Scoring table: BLOSUM62
 Gapop 10.0 , Gapext 0.5

Searched: 2443163 seqs, 439378781 residues

Total number of hits satisfying chosen parameters: 146

Minimum DB seq length: 0
 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 100%
 Listing first 500 summaries

Database : A_Geneseq_21:
 1: geneseq201980s:
 2: geneseq201990s:
 3: geneseq20000s:
 4: geneseq20010s:
 5: geneseq20020s:
 6: geneseq20030s:
 7: geneseq2003bs:
 8: geneseq2005as:
 9: geneseq2005as:
 9:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query	Match Length	DB ID	Description
1	846	100.0	165	3 AAY33445	Aay93445 Amino aci
2	846	100.0	165	3 AAB03760	Aab03760 Human ery
3	846	100.0	165	3 AAY94605	Aay94605 Human ery
4	846	100.0	165	3 AAY9705	Aay99705 Non-glyco
5	846	100.0	165	4 AAB94525	Aab94525 Amino aci
6	846	100.0	165	4 ABB93621	Abb93621 Protein #
7	846	100.0	165	4 ABB6697	Abb6697 Human ery
8	846	100.0	165	5 AAM23061	Aam53061 Human ery
9	846	100.0	165	5 ABT77896	Abt77896 Amino aci
10	846	100.0	165	6 ABP98492	Abp98492 Amino aci
11	846	100.0	165	6 ABR39995	Ab39995 Human ery
12	846	100.0	165	8 ADI06780	Adl06780 Human 165
13	846	100.0	165	8 ADN9745	Adn9745 Mature hu
14	846	100.0	165	8 AD059415	Ad059415 Human 165
15	846	100.0	165	8 ADU74421	Adu74421 Mature hu
16	846	100.0	165	9 AEK7164	Aek7164 ERYthropo
17	846	100.0	165	9 AEB21317	Aeb21317 Amino aci
18	846	100.0	165	1 AAPT0398	Aap0398 Sequence
19	846	100.0	166	2 AAR23593	Aar23593 Recombina
20	846	100.0	166	2 AAW88404	AAw88404 Human ery
21	846	100.0	166	2 AAW7780	AAw7780 Human EPO
22	846	100.0	166	3 ABB07030	Abb07030 Modified
23	846	100.0	166	4 AAB83622	Aab83622 Protein #
24	846	100.0	166	4 AAE02641	Aae02641 Human ery
201	9				AEC05278 Modified

Abb6698 Human ery

Abb92101 Human ery

Aam53062 Human ery

Abt77897 Amino aci

Adg65661 Human ery

Abr3996 Human ery

Abz57500 Human ery

Adf70339 Human ery

Adl92150 Human ery

Abk70564 Human ery

Adl8867 Human cyt

Adl06781 Human 166

Ado59416 Human 166

Adv67303 Human aci

Ady93798 Human ery

Aea47165 ERYthro

Aeb21318 Amino aci

Aap50299 Human rec

Aap50298 Human rec

Abb77899 Amino aci

Abb77898 Amino aci

Abp0300 Human alb

Adf16727 Human alb

Aep60597 Clone lam

Aap1195 ERYthro

Adf1588 Human alb

Adf1589 Human alb

Adf16589 Human alb

Adf16590 Human alb

Adf16727 Human alb

Adf16726 Human alb

Adf15298 Human alb

Adf15295 Human alb

Adf16587 Human alb

Adf16588 Human alb

Adf16589 Human alb

Adf16590 Human alb

Adf16727 Human alb

Adf16726 Human alb

Adf16728 Human alb

Adf16589 Human alb

Adf16588 Human alb

Adf16587 Human alb

Adf16588 Human alb

98 846 100.0 205 8 ADJ71846 XX
 99 846 100.0 209 7 ADJ79063 PR 06-NOV-1998; 9BAR-00105609.
 100 846 100.0 220 5 ABB7939 PR 23-FEB-1999; 9BAR-00100679.
 101 846 100.0 220 7 ABR57656 XX
 102 846 100.0 302 2 AAR23596 PA (STER-) STERRENBEELD BIOTECHNOLOGIE NORTH AMERICA.
 103 846 100.0 303 2 AAR23598 XX
 104 846 100.0 321 2 AAR23075 PI Caccagno CM, Criscuolo M, Melo C, Vidal JA;
 105 846 100.0 321 2 AAR23597 XX
 106 846 100.0 330 2 AAR23076 DR WPI; 2000-376574/32.
 107 846 100.0 340 2 AAR23078 PT New host cell producing recombinant human erythropoietin (EPO) used for
 108 846 100.0 349 2 AAR23079 PT large scale production of EPO.
 109 846 100.0 370 7 ADO7962 XX
 110 846 100.0 376 2 AAW99360 PS Claim 1; Page 26-27; 51pp; English.
 111 846 100.0 379 9 AEB12283 XX
 112 846 100.0 428 7 ABU64200 CC The present sequence represents human erythropoietin protein. The specification describes a host cell line which is used to produce human erythropoietin (EPO). EPO is a glycoprotein. The cell line is used for the production of recombinant human erythropoietin. The protein is used for the treatment of anaemia, especially anaemia derived from renal failure.
 113 846 100.0 428 8 ADD010513 CC
 114 846 100.0 428 9 ADV97050 CC
 115 846 100.0 435 7 ADM33857 CC
 116 846 100.0 435 8 ADR48888 CC
 117 846 100.0 435 8 ADW47520 CC
 118 846 100.0 435 9 AEA18937 CC
 119 846 100.0 435 9 AEA88757 XX
 120 846 100.0 435 9 AEA88757 SQ Sequence 165 AA:
 121 846 100.0 436 7 ADD33853 Query Match 100.0%; Score 846; DB 3; Length 165;
 122 846 100.0 436 8 ADR48884 Best Local Similarity 100.0%; Pred. No. 2.2e-86;
 123 846 100.0 436 8 AEW47516 Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;
 124 846 100.0 436 9 AEA18933 Human ery XX
 125 846 100.0 436 9 AEA88753 Human ery XX
 126 846 100.0 437 7 ADM33855 Human Hub XX
 127 846 100.0 437 8 ADR48886 HubEO-L-V XX
 128 846 100.0 437 9 AEW47518 Human EPO Db 1 APPRLICDSRVLEERYLLEAKRAENITTGCAEHCSLNENITVPTKVNHYAWKMEV63QAA 60
 129 846 100.0 437 9 AEA18935 Human ery QY 61 VEVMOGLALISEAVYLRGQALVNSSQPWPLOQEVDDKAVSGLSLTTLRAGQKEAIS 120
 130 846 100.0 437 9 AEA88755 Human ery QY 61 VEVMOGLALISEAVYLRGQALVNSSQPWPLOQEVDDKAVSGLSLTTLRAGQKEAIS 120
 131 846 100.0 768 7 ADF16425 Human alb Db Adf16425 Human alb 121 PPDASAAPERTTADTEPKLFRYTSNFGRKLUKYTBACRGD 165
 132 846 100.0 768 7 ADF16426 Human alb QY Adf16426 Human alb 121 PPDASAAPLRTTADTEPKLFRVYSNFLRGKLUKYTBACRGD 165
 133 846 100.0 768 7 ADF16424 Human alb Adf16424 Human alb
 134 846 100.0 768 7 ADF16424 Human alb Adf16424 Human alb
 135 846 100.0 768 7 ADF15091 Human alb Adf16563 Human alb
 136 846 100.0 768 7 ADF15091 Human alb Adf16563 Human alb
 137 846 100.0 769 7 ADF15091 Human alb Adf15091 Human alb
 138 846 100.0 777 7 ADF15082 Human alb Adf15082 Human alb
 139 846 100.0 777 7 ADF15082 Human alb Adf15078 Human alb
 140 846 100.0 777 7 ADF15075 Human alb Adf15075 Human alb
 141 846 100.0 777 7 ADF15079 Human alb Adf15079 Human alb
 142 846 100.0 777 7 ADF15081 Human alb Adf15081 Human alb
 143 846 100.0 777 7 ADF15113 Human alb Adf15113 Human alb
 144 846 100.0 951 7 ADF15113 Human alb Adf15113 Human alb
 145 846 100.0 951 7 ADF15108 Human alb Adf15108 Human alb
 146 846 100.0 954 7 ADF15105 Human alb Adf15105 Human alb

ALIGNMENTS

RESULT 1
 AAY33445 ID AAY33445 standard; protein; 165 AA.
 XX AC AAY33445;
 XX DT 04-SEP-2000 (first entry)
 DE Amino acid sequence of human erythropoietin.
 KW Human; erythropoietin; EPO; anaemia; renal failure.
 OS Homo sapiens.
 XX PI Carcagno CM, Criscuolo M, Melo C, Vidal JA;
 XX PN WO200028066-A1.
 XX PR 06-NOV-1998; 9BAR-00105611.
 XX PR 23-FEB-1999; 9BAR-00100681.
 XX PR (STER-) STERRENBEELD BIOTECHNOLOGIE NORTH AMERICA.
 XX PI Carcagno CM, Criscuolo M, Melo C, Vidal JA;
 XX DR WPI; 2000-376519/32.
 XX PT A novel method for the massive culture of recombinant mammalian cells producing recombinant human erythropoietin.

PP 08-NOV-1999; 99WO-US026238.

XX
XX
PS Example 8; Page 11-12; 23pp; English.

CC This sequence represents the human erythropoietin amino acid sequence.

CC Erythropoietin is a glycoprotein that stimulates erythroblast CC differentiation in the bone marrow. The present invention relates to a CC method for the large scale production of human EPO from recombinant CC mammalian cells. The method comprises culturing mammalian cells which CC express recombinant human EPO in culture medium comprising insulin.

CC Erythropoietin can be used to treat anaemia derived from renal failure. CC The method allows for the industrial scale production of EPO, and CC overcomes the problems of low reproducibility and output quality which CC are encountered with previous production methods

XX SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 3; Length 165;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEARENITTCGCAEBCSLENINITYPDTKVNFYAWRMEVCGQA 60
Db 1 APPRLICDSRVLERYLLEAKEARENITTCGCAEBCSLENINITYPDTKVNFYAWRMEVCGQA 60
Qy 61 VEWQGALLSSEAVLRGQALLVNSQPEPLQLHDKAVSGRSLTTRALGAQKEAIS 120
Db 61 VEWQGALLSSEAVLRGQALLVNSQPEPLQLHDKAVSGRSLTTRALGAQKEAIS 120
Qy 121 PPDASAAAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165
Db 121 PPDASAAAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165

RESULT 3
AY94605

ID AAY94605 standard; protein; 165 AA.

XX AC AAY94605;

XX DT 28-NOV-2000 (first entry)

XX DE Human erythropoietin.

XX KW Human; erythropoietin; EPO; purification; anaemia.

XX OS Homo sapiens.

XX FR Key Location/Qualifiers

FT Modified-site 24 /note= "N-Glycosylation site"

FT Modified-site 38 /note= "N-Glycosylation site"

FT Modified-site 83 /note= "N-Glycosylation site"

FT Modified-site 126 /note= "O-Glycosylation site"
/note= "O-Glycosylation site"

XX WO200027869-A1.

XX PD 18-MAY-2000.

XX PP 08-NOV-1999; 99WO-US026241.

XX PR 06-NOV-1998; 98AR-00105610.
PR 23-FEB-1999; 99AR-00100680.

XX PA (STER-) STERRENBELD BIOTECHNOLOGIE NORTH AMERICA.
XX PI Carcagno CM, Criscuolo M, Melo C, Vidal JA;
XX DR WPI; 2000-376485/32.

XX PT Novel methods for purifying recombinant human erythropoietin from

PT mammalian cell culture reagents.

XX PS Claim 16; Page 18; 30pp; English.

CC The present invention relates to a method for purifying erythropoietin (EPO) for treatment of disease, especially anaemia. The method involves treating cell culture supernatants with differential precipitation, hydrophobic interaction chromatography, diafiltration, anionic and cationic exchange chromatography and molecular exclusion chromatography.

CC The present sequence is the protein from the culture supernatant of transfected cell lines, after purification by the above process. The sequence shows total homology with natural human EPO. The advantage of this method is that high purity and quality EPO is produced. A further advantage is that the process does not involve the use of organic solvents that may harm the environment

XX SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 3; Length 165;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEARENITTCGCAEBCSLENINITYPDTKVNFYAWRMEVCGQA 60
Db 1 APPRLICDSRVLERYLLEAKEARENITTCGCAEBCSLENINITYPDTKVNFYAWRMEVCGQA 60
Qy 61 VEWQGALLSSEAVLRGQALLVNSQPEPLQLHDKAVSGRSLTTRALGAQKEAIS 120
Db 61 VEWQGALLSSEAVLRGQALLVNSQPEPLQLHDKAVSGRSLTTRALGAQKEAIS 120
Qy 121 PPDASAAAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165
Db 121 PPDASAAAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165

RESULT 4
AY99705

ID AY99705 standard; protein; 165 AA.

XX AC AAY99705;

XX DT 15-SEP-2000 (first entry)

XX DE Non-glycosylated erythropoietin analogue NGE-166delta.

XX KW Human; non-glycosylated erythropoietin analogue; NGEA; haematocrit; antianaemic; anaemia; erythropoiesis promoter; mutant; mutein.

XX OS Homo sapiens.

XX OS Synthetic.

XX PN WO20032772-A2.

XX PD 08-JUN-2000.

XX PP 23-NOV-1999; 99WO-US027801.

XX PR 30-NOV-1998; 98US-0110289P.

XX PA (BLIL) LILLY & CO ELL.

XX PI Beals JM, Glaesner W, Micanovic R, Millican RL, Witcher DR;

XX DR WPI; 2000-412320/35.

XX DR N-PSDB; AAA48373.

XX PT

Non-glycosylated erythropoietin compound useful for increasing hematocrit level in mammal with insufficient hematocrit levels in conditions such as anemia, comprises protein covalently bonded to polymer.

XX PS

Claim 2; Page 93-94; 94pp; English.

XX

The present sequence is a non-glycosylated erythropoietin analogue (NGEA)

CC designated NGB-166delta. The protein sequence is identical to the
 CC sequence of wild-type human non-glycosylated erythropoietin NGB except
 CC that Arg at position 165 is deleted. NGB promotes erythropoiesis and can
 CC therefore be used to increase haematocrit levels in mammals with
 CC conditions such as anaemia, in which levels of haematocrit are
 CC insufficient. NGB analogues can also be used to treat such conditions.
 CC NGBs do not themselves cause a significant increase in haematocrit but
 CC they acquire that property once they are derivatised with polyethylene
 CC glycol polymers. The analogues can be produced using a linkerless
 CC aldehyde modification process. They show stability and bioactivity in
 CC vivo. The nucleotide sequence encoding this protein was constructed
 CC synthetically by in vitro hybridisation using a set of six overlapping
 CC oligonucleotides from the positive strand of human erythropoietin cDNA
 CC with six complementary oligonucleotides (negative strand). The codon
 CC usage was 100% optimised for E. coli codon usage. The hybridised
 CC oligonucleotides were ligated with T4 DNA ligase and the ligation product
 CC amplified by PCR. The nucleotide sequence was used to express the protein
 XX in host cells.

SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 3; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2 2e-86;
 CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKENAINTTGCAGHCSLNENITVDPDTKUNFIAWKRMEVGQAA 60
 Db 1 APPRLICDSRVRLERYLLEAKENAINTTGCAGHCSLNENITVDPDTKUNFIAWKRMEVGQAA 60

QY 61 VEWMOGLALISEAVLRGQALLVNSQWPQLHQVADKAVSGRSLSLTLLRAGQKEAIS 120
 Db 61 VEWMOGLALISEAVLRGQALLVNSQWPQLHQVADKAVSGRSLSLTLLRAGQKEAIS 120

QY 121 PPDASAAPIRITTADETRKLFRVYSNFRGKLYTGEARCTGD 165
 Db 121 PPDASAAPIRITTADETRKLFRVYSNFRGKLYTGEARCTGD 165

RESULT 5

AAB84525 AAB84525 standard; protein; 165 AA.

ID AAB84525;

XX AC AAB84525;

XX DT 05-SEP-2001 (first entry)

XX DE Amino acid sequence of human erythropoietin (EPO) protein.

XX KW Erythropoietin; EPO; erythropoietin stimulating protein; NESP;
 KW sustained release.

XX OS Homo sapiens.

XX PN WO201303220-A1.

XX PD 03-MAY-2001.

XX PP 23-OCT-2000; 2000W0-US0229257.

XX PR 22-OCT-1999; 99US-00426566.
 13-OCT-2000; 2000US-00687981.
 (AMGE-) AMGEN INC.

PT Burke P, Klumb L, Murphy K, Herberger J, French DJ;

XX DR WPI; 2001-417552/44.

XX Sustained release composition comprises an active biological ingredient,
 PT notably a protein or other biopolymer, particularly erythropoietin
 PT stimulating protein, in biocompatible, biodegradable polymeric
 PT microparticles.

SQ Sequence 165 AA;

PS Disclosure; Page 56; 61pp; English.

XX The present sequence encodes a human erythropoietin (EPO) protein. The
 CC specification describes a composition for the sustained release of
 CC biologically active EPO stimulating protein (NESP). The reduced frequency
 CC of administration of NESP, which requires preferably injection by skilled
 CC personnel, improves patient compliance. Also, sustained release reduces
 CC the nature and severity of any side effects of the drug

SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 4; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2 2e-86;
 CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKENAINTTGCAGHCSLNENITVDPDTKUNFIAWKRMEVGQAA 60
 Db 1 APPRLICDSRVRLERYLLEAKENAINTTGCAGHCSLNENITVDPDTKUNFIAWKRMEVGQAA 60

QY 61 VEWMOGLALISEAVLRGQALLVNSQWPQLHQVADKAVSGRSLSLTLLRAGQKEAIS 120
 Db 61 VEWMOGLALISEAVLRGQALLVNSQWPQLHQVADKAVSGRSLSLTLLRAGQKEAIS 120

RESULT 6

ABB83621 ABB83621 standard; protein; 165 AA.

XX AC ABB83621;

XX DT 10-OCT-2002 (first entry)

XX DB Protein #1 relating to modified erythropoietin glycoprotein.

XX KW Erythropoietin glycoprotein; anaemia; chronic renal failure; AIDS;
 KW cancer.

XX OS Unidentified.

XX PN NO200003372-A.

XX PD 03-JAN-2001.

XX PF 28-JUN-2000; 2000ONO-00003372.

XX PR 02-JUL-1999; 99US-0142254P.
 PR 23-AUG-1999; 99US-0150225P.
 PR 31-AUG-1999; 99US-0151548P.
 PR 17-NOV-1999; 99US-0166151P.

XX PA (Hoffmann La Roche & Co AG F.

XX PI Bailon PS;

XX DR WPI; 2001-135308/14.

XX PT New conjugate having modified erythropoietin glycoprotein useful for
 PT stimulating red blood cell production and for treating diseases
 PT correlated with anemia in chronic renal failure, AIDS or cancer patients.

XX Disclosure; Page 21-22; 30pp; Norwegian.

XX CC This invention relates to new conjugate having a modified erythropoietin
 CC glycoprotein, useful for stimulating red blood cell production, and for
 CC treating or preventing diseases correlated with anaemia in chronic renal
 CC failure, AIDS or cancer patients. The present sequence is a protein
 CC related to the invention

SQ Sequence 165 AA;

CC immunodeficiency syndrome), and/or for the treatment of cancer patients undergoing chemotherapy. Unlike prior art erythropoietin compositions, the compositions of the invention do not contain human serum albumin (thereby avoiding the possibility of viral infections and allergic reactions associated with this component), are liquid rather than lyophilisates (and therefore do not need to be reconstituted before administration), and are stable at elevated temperatures such as 25 degrees Celsius and even 40 degrees Celsius, and therefore can be stored without refrigeration for prolonged periods without degradation and loss of activity. The present sequence represents the 165 residue form of human erythropoietin which is specifically claimed for use in a composition of the invention.

CC Sequence 165 AA;

Query Match 100.0%; Score 846; DB 5; Length 165;
Best Local Similarity 100.0%; Pred. No. 2-2e-86; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLYRVLLEKAENITTCGAECHCSLENNTIVPDTKVNFKWAKMEVGQAA 60
Db 1 APPRLICDSRVLYRVLLEKAENITTCGAECHCSLENNTIVPDTKVNFKWAKMEVGQAA 60

QY 61 VEWMOGLALISEAVLRLQGALLVNSSQPMPLQLHVDAVKGSLRSLTLLRALGAQEAIIS 120
Db 61 VEWMOGLALISEAVLRLQGALLVNSSQPMPLQLHVDAVKGSLRSLTLLRALGAQEAIIS 120

QY 121 PPDAAASAPRITTADETRKLFRVYSNFLRGKLUYTGEACRTGD 165
Db 121 PPDAAASAPRITTADETRKLFRVYSNFLRGKLUYTGEACRTGD 165

RESULT 9

ID ABB77896
XX ABB77896 standard; protein; 165 AA.

AC XX
XX 07-OCT-2002 (first entry)
DB Amino acid sequence of a human erythropoietin (EPO).

XX Human; erythropoietin; EPO; glycoprotein; reticulocyte production; red blood cell production; anaemia; chronic renal failure; acquired immunodeficiency syndrome; AIDS; cancer; bone marrow; committed erythroid progenitor.
XX Homo sapiens.
XX WO200249673-A2.
XX PN
XX PD 27-JUN-2002.
XX PR 08-DEC-2001; 2001WO-EP014434.
XX 20-DEC-2000; 2000EP-00127891.
XX PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX Burg J, Engel A, Franze R, Hilger B, Schuring HE, Tischer W; PI Wozny M; DR XX WPI; 2002-566640/60.
XX Novel conjugate of erythropoietin glycoprotein with polyethylene glycol, useful for treating diseases correlated with anaemia in chronic renal failure patients and acquired immunodeficiency syndrome.

XX Claim 26; Fig 1; 40pp; English.
PS The present sequence represents a human erythropoietin (EPO) protein. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal

CC alpha-amino group, chosen from human EPO (hEPO) or its analogues where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increases circulating half-life and plasma residence time, decreased clearance, increased clinical activity in vivo, improved potency and stability, when compared to unmodified EPO. The EPO conjugate is useful for preparing medications for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

XX Sequence 165 AA;

Query Match 100.0%; Score 846; DB 5; Length 165;
Best Local Similarity 100.0%; Pred. No. 2-2e-86; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLYRVLLEKAENITTCGAECHCSLENNTIVPDTKVNFKWAKMEVGQAA 60
Db 1 APPRLICDSRVLYRVLLEKAENITTCGAECHCSLENNTIVPDTKVNFKWAKMEVGQAA 60

QY 61 VEWMOGLALISEAVLRLQGALLVNSSQPMPLQLHVDAVKGSLRSLTLLRALGAQEAIIS 120
Db 61 VEWMOGLALISEAVLRLQGALLVNSSQPMPLQLHVDAVKGSLRSLTLLRALGAQEAIIS 120

QY 121 PPDAAASAPRITTADETRKLFRVYSNFLRGKLUYTGEACRTGD 165
Db 121 PPDAAASAPRITTADETRKLFRVYSNFLRGKLUYTGEACRTGD 165

RESULT 10

ID ABP98492
XX ABP98492 standard; protein; 165 AA.

AC XX
XX DT 29-JUL-2003 (first entry)
DB Amino acid sequence of human erythropoietin (EPO).

XX Human; erythropoietin; EPO; novel erythropoiesis stimulating protein; KW NESP; haemocrit level.
XX OS Homo sapiens.
XX WO2003020299-A1.
XX PN
XX PD 13-MAR-2003.
XX PR 29-AUG-2002; 2002WO-US027855.
XX 30-AUG-2001; 2001US-0094517.
XX PA (KIRIN) KIRIN AMGEN INC.
XX PI Li T, Chang BS, Sloey C;
XX DR WPI; 2003-402847/38.
XX PT Pharmaceutical formulation for single use comprises biologically active agent, methionine and optional preservative and does not contain human serum albumin.

PS Claim 6; Page 37; 40PP; English.
XX The present sequence represents human erythropoietin (EPO). EPO is used as the active agent in formulations of the invention. The specification describes a pharmaceutical formulation, which comprises a biologically active agent (e.g. EPO or novel erythropoiesis stimulating protein

CC (NESP), methionine and a preservative. The formulation does not contain
 CC human serum albumin (HSA). The formulation has improved stability.
 CC Incorporation of methionine and other stabilizing agents into the
 CC formulation produces a more stable formulation, even in extreme
 CC conditions, where the critical degradations induced by light, heat,
 CC impurities in additives, leachates in the prefilled syringes, the
 CC manufacturing process, storage, transportation and handling are
 CC prevented. The formulation is useful as a single use and a multi-dose
 CC formulation. Where NESP is the active agent, it may be used to raise
 XX haemocrit levels
 SQ sequence 165 AA;

Query Match 100.0%; Score 846; DB 6; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2.2e-66; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; MisMatches 0; Del 0; Insert 0;

QY 1 APPRLICDSRVLERYLRAKEARNITTCARICSLNENITVPTPKVNYANTRMEVQQA 60
 1 APPRLICDSRVLERYLRAKEARNITTCARICSLNENITVPTPKVNYANTRMEVQQA 60
 Db 61 VEWMOGLALLSEAVLRGQALLVNSQWEPLQLHVDKAVSGLSLTLLRALGAOKEAIIS 120
 QY 121 PPDAAASAPLRTTADTPRKLFRVYSNPLRGKLYTGACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFRVYSNPLRGKLYTGACRTGD 165

RESULT 11

ABR3995

ID ABR3995 standard; protein; 165 AA.

AC ABR3995;

XX DT 02-SEP-2003 (first entry)

XX DE Human erythropoietin (EPO) sequence.

XX KW EPO; erythropoietin; mitein; reticulocyte; red blood cell; antianemic;

XX AIDS; cancer;

XX OS Homo sapiens.

XX FH Key Location/Qualifiers

FT Disulfide-bond 7..161 /note= "disulphide bridge"

FT Disulide-bond 29..33 /note= "disulphide bridge"

FT Modified-site 38 /note= "Asn is N-glycosylated"

FT Modified-site 83 /note= "Asn is N-glycosylated"

FT Modified-site 126 /note= "Ser is O-glycosylated"

/note= "Ser is O-glycosylated"

XX WO2003029291-A2.

XX PD 10-APR-2003.

XX 20-SEP-2002; 2002WO-EP010556.

XX 25-SEP-2001; 2001EP-00122555.

XX PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX PT Tischer W;

XX DR WPI; 2003-457226/43.

PT Novel erythropoietin mitein having in vivo biological activity of causing
 CC bone marrow cells to increase production of reticulocytes/red blood

PT cells, is N-glycosylated at Asn38 and Asn83 but not N-glycosylated at
 PT Asn24. The invention relates to an erythropoietin mitein (I) having the in vivo
 XX Claim 6; Page 21-22; 22pp; English.
 XX The invention relates to an erythropoietin mitein (I) having the in vivo
 CC biological activity of causing bone marrow cells to increase production
 CC of reticulocytes and red blood cells, characterized by being N-
 CC glycosylated at Asn38 and Asn83 but not N-glycosylated at Asn24. (I) or
 CC an aqueous composition comprising an erythropoietin mitein is useful for
 CC the preparation of a medicament for the treatment or prophylaxis of
 CC diseases correlated with anemia in chronic renal failure patients (CRF),
 CC AIDS and for the treatment of cancer patients undergoing chemotherapy.
 CC (I) or the composition is useful for treating a human patient
 CC experiencing blood disorders characterized by low or defective red blood
 CC cell production. (I) is useful for enhancing red blood cell formation.
 CC The present sequence represents a human erythropoietin (EPO) sequence
 XX SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 6; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2.2e-66; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; MisMatches 0; Del 0; Insert 0;

QY 1 APPRLICDSRVLERYLRAKEARNITTCARICSLNENITVPTPKVNYANTRMEVQQA 60
 1 APPRLICDSRVLERYLRAKEARNITTCARICSLNENITVPTPKVNYANTRMEVQQA 60
 Db 61 VEWMOGLALLSEAVLRGQALLVNSQWEPLQLHVDKAVSGLSLTLLRALGAOKEAIIS 120
 QY 121 PPDAAASAPLRTTADTPRKLFRVYSNPLRGKLYTGACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFRVYSNPLRGKLYTGACRTGD 165

RESULT 12

ADL06780

ID ADL06780 standard; protein; 165 AA.

XX AC ADL06780;

XX DT 03-JUN-2004 (first entry)

XX DE Human 165 residue erythropoietin (EPO), SEQ ID NO:1.

XX Human; erythropoietin; EPO; iron distribution disturbance; diabetes;

XX non-insulin dependent diabetes; type 2 diabetes; reticulocyte production;

XX red blood cell production; antidiabetic.

XX OS Homo sapiens.

XX RN WO2004019972-A1.

XX PD 11-MAR-2004.

XX PF 20-AUG-2003; 2003WO-EP009194.

XX PR 29-AUG-2002; 2002EP-00019100.

XX (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX PI Lehmann P, Roeddiger R, Walter-Matsui R;

XX DR WPI; 2004-282643/26.

XX PT Use of erythropoietin protein in manufacture of medicament for treating

XX PT disturbances of iron distribution in diabetes.

XX PS Claim 6; SEQ ID NO 1; 31pp; English.

XX The invention relates to the use of an erythropoietin (EPO) protein for

the treatment of disturbances of iron distribution in diabetes. The erythropoietin protein is preferably a human erythropoietin (e.g., epoetin alpha and epoetin beta) which may be expressed by endogenous gene activation or an erythropoietin analogue such as darbepoietin alpha. The erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation. Patients with diabetes have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effects of iron accumulation in certain organs, leading to organ damage and destruction, and/or experience effects similar to anaemia due to iron usage in blood cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticularocytes and red blood cells, and this has been found to have a beneficial effect on iron distribution disturbances in diabetes e.g., non-insulin dependent (type 2) diabetes. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron distribution in diabetes. The present sequence represents a 165 amino acid human erythropoietin which is specifically claimed for use in the invention.

XX
SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 8; Length 165;
Best Local Similarity 100.0%; Pred. No. 2-2e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKENAINTTCGAEHCSLENNTIVPTKVNHYAWKMEVGQAA 60
Db 1 APPRLICDSVRLERYLLEAKENAINTTCGAEHCSLENNTIVPTKVNHYAWKMEVGQAA 60
QY 61 VEWVOGLALLSEAVLRGQALLVNSQPWPQLQHVDAVSGLSLTTLRGAQKEAIS 120
Db 61 VEWVOGLALLSEAVLRGQALLVNSQPWPQLQHVDAVSGLSLTTLRGAQKEAIS 120
QY 121 PPDAAASAPLRTTADTRKLFRVYSNPLGKLYTGACRTGD 165
Db 121 PPDAAASAPLRTTADTRKLFRVYSNPLGKLYTGACRTGD 165

RESULT 13

ADN49745
ID ADN49745 standard; protein; 165 AA.
AC XX
XX
AC ADN49745;
DT 15-JUL-2004 (first entry)

Mature human erythropoietin protein SeqID 73.

human; erythropoietin; EPO; glycoconjugation; glycopEgylated EPO peptide; anaemia; antianaemic; haemocrit level; kidney dialysis; haematology; erythropoietin.

XX
OS Homo sapiens.
XX
WO2004033651-A2.
XX
PN 22-APR-2004.
XX
PP 08-OCT-2003; 2003WO-US031974.

XX
PR 09-OCT-2002; 2002WO-US032263.

PR 05-NOV-2002; 2002US-00287994.

PR 06-JAN-2003; 2003US-00360770.

PR 19-FEB-2003; 2003US-00360779.

PR 09-APR-2003; 2003US-00410945.

XX
PA (NEOS-) NEOSE TECHNOLOGIES INC.

PI De Frees S, zopf D, Bayer R, Bowe C, Hakes D, Chen X;

DR XX
WPI; 2004-399848/37.

XX
PT Novel erythropoietin Peptide comprising one or more glycans, having PT glycoconjugate molecule covalently attached to peptide, useful for PT treating anemia in mammal such as human.

XX
PS Claim 38; SEQ ID NO 73; 1018PP; English.

XX
CC This invention relates to novel erythropoietin (EPO) peptides and the CC remodelling and glycoconjugation of these naturally occurring peptides CC thereof. Specifically, each EPO peptide comprises one or more glycans and CC has a glycoconjugate molecule such as polyethylene glycol (PEG) attached CC to it. Accordingly, the present invention provides glycopEGylated EPO peptides that have either monoantennary, biantennary or triantennary CC glycans covalently attached thereto. As such, these peptides are useful CC for the treatment of anaemia, and hence exhibit anti-anaemic activities CC working to increase haemocrit levels in mammals, in particular in humans i.e. increasing the relative volume of blood occupied by CC erythrocytes. Furthermore, EPO therapy can be used to treat kidney CC dialysis patients. This polypeptide is a human protein sequence related CC to the field of haematology, given in an exemplification of the invention.

XX
SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 8; Length 165;
Best Local Similarity 100.0%; Pred. No. 2-2e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKENAINTTCGAEHCSLENNTIVPTKVNHYAWKMEVGQAA 60
Db 1 APPRLICDSVRLERYLLEAKENAINTTCGAEHCSLENNTIVPTKVNHYAWKMEVGQAA 60
QY 61 VEWVOGLALLSEAVLRGQALLVNSQPWPQLQHVDAVSGLSLTTLRGAQKEAIS 120
Db 61 VEWVOGLALLSEAVLRGQALLVNSQPWPQLQHVDAVSGLSLTTLRGAQKEAIS 120
QY 121 PPDAAASAPLRTTADTRKLFRVYSNPLGKLYTGACRTGD 165
Db 121 PPDAAASAPLRTTADTRKLFRVYSNPLGKLYTGACRTGD 165

RESULT 14

ID AD059415
ID AD059415 standard; protein; 165 AA.
AC XX
AC AD059415;
DT 26-AUG-2004 (first entry)

XX
DR Human 165 residue erythropoietin (EPO), SEQ ID NO:1.
XX
Human; erythropoietin; EPO; iron distribution disturbance; heart disease; KW heart insufficiency; coronary heart disease; atherosclerosis; acute coronary syndrome; heart failure; congestive heart failure; KW reticulocyte production; red blood cell production; cardiotonic; KW anti-arteriosclerotic.
XX
OS Homo sapiens.
XX
PN WO2004047858-Al.

XX
PP 10-JUN-2004.
XX
PP 17-NOV-2003; 2003WO-EP012822.
XX
PR 22-NOV-2002; 2002EP-00026342.
XX
PA (HOFF) HOFFMANN LA ROCHE & CO AG F.
XX
PI Lehmann P, Roediger R, Walter-Matsui R;
XX
DR WPI; 2004-450212/42.

PT Use of erythropoietin protein in the manufacture of medicament for
 PT treating disturbances of iron distribution in heart diseases e.g. heart
 PT insufficiency.
 XX
 PS Claim 6; SEQ ID NO 1; 31PP; English.

The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in heart diseases. The erythropoietin protein is preferably a human erythropoietin (e.g., epoetin alpha and epoetin beta) which may be expressed by endogenous gene activation or an erythropoietin analogue such as darbepoietin alpha. The erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation. Patients with heart diseases have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effects of iron accumulation in certain organs, leading to organ damage and destruction, and/or experience effects similar to anaemia due to iron usage in blood cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticulocytes and red blood cells, and this has been found to have a beneficial effect on iron distribution disturbances in heart diseases e.g., heart insufficiency, coronary heart disease, atherosclerosis, acute coronary syndrome, heart failure and congestive heart failure. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron distribution in heart diseases. The present sequence represents a 165 amino acid human erythropoietin which is specifically claimed for use in the invention.

CC Sequence 165 AA;
 SQ

Query Match 100.0%; Score 846; DB 8; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLYRVLRAKEAENITTCGCAECSLNEINTVPTDKVNYAWKRMVEVGQA 60
 Db 1 APPRLICDSRVLYRVLRAKEAENITTCGCAECSLNEINTVPTDKVNYAWKRMVEVGQA 60
 QY 61 VEWMOGLALLSEAVLRQGQALLVNSQSPWPLQLAVDKAVSGRSLSLTILRAIGAQEAIS 120
 Db 61 VEWMOGLALLSEAVLRQGQALLVNSQSPWPLQLAVDKAVSGRSLSLTILRAIGAQEAIS 120
 QY 121 PPDASAAPLRTTADTRKLFRVYSNTRGKLUKYGEACTGD 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNTRGKLUKYGEACTGD 165

SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 8; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

AC ADU74421;
 XX DT 10-FEB-2005 (first entry)
 DE Mature human erythropoietin.
 KW Hemostatic; Hepatotropic; Antianemic; Cytostatic; Osteopathic; Antibacterial; Respiratory-Gen.; Antiinflammatory; Nephrotopic; Antifertility; Antitubercular; Tuberculosis; protein engineering; bleeding; factor IX deficiency; liver cirrhosis; infertility; anemia; end-stage renal disease; acute myelogenous leukemia; osteoporosis; pulmonary fibrosis; tuberculosis; cryptococcal meningitis; etc. The glycopptide produced using (M1) has specific customized or desired glycosylation pattern. (M1) allows efficient production of a protein remodelled in the present invention

SQ Sequence 165 AA;

Query Match 100.0%; Score 846; DB 8; Length 165;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLYRVLRAKEAENITTCGCAECSLNEINTVPTDKVNYAWKRMVEVGQA 60
 Db 1 APPRLICDSRVLYRVLRAKEAENITTCGCAECSLNEINTVPTDKVNYAWKRMVEVGQA 60
 QY 61 VEWMOGLALLSEAVLRQGQALLVNSQSPWPLQLAVDKAVSGRSLSLTILRAIGAQEAIS 120
 Db 61 VEWMOGLALLSEAVLRQGQALLVNSQSPWPLQLAVDKAVSGRSLSLTILRAIGAQEAIS 120
 QY 121 PPDASAAPLRTTADTRKLFRVYSNTRGKLUKYGEACTGD 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNTRGKLUKYGEACTGD 165

RESULT 15
 ADU74421
 ID ADU74421 standard; protein; 165 AA.
 XX
 AC ADU74421;
 XX DT 10-FEB-2005 (first entry)
 DE Mature human erythropoietin.
 KW Hemostatic; Hepatotropic; Antianemic; Cytostatic; Osteopathic; Antibacterial; Respiratory-Gen.; Antiinflammatory; Nephrotopic; Antifertility; Antitubercular; Tuberculosis; protein engineering; bleeding; factor IX deficiency; liver cirrhosis; infertility; anemia; end-stage renal disease; acute myelogenous leukemia; osteoporosis; pulmonary fibrosis; tuberculosis; etc. The glycopptide produced using (M1) has specific customized or desired glycosylation pattern. (M1) allows efficient production of a protein remodelled in the present invention

RESULT 16
 AEA47164
 ID AEA47164 standard; protein; 165 AA.
 XX
 AC AEA47164;
 XX DT 11-AUG-2005 (first entry)

RESULT 18
 ID AAP70398
 DE standard; protein; 166 AA.
 XX
 AC AAP70398;
 XX
 DT 19-FEB-1991 (first entry)
 DE Sequence of human erythropoietin (EPO).
 KW Megakaryocyte-platelet growth factor; hormone;
 KW megakaryocyte colony stimulating factor; therapy;
 KW small acetyl cholinesterase positive cell; erythrocyte growth effect.
 OS Homo sapiens.
 XX
 PN JPF2149624-A.
 XX
 PR 13-SEP-1985; 85UP-00203049.
 XX
 PA (ORTH) ORTHO PHARM CORP.
 XX
 PT Rosen JT;
 XX
 DR WPI; 1992-150819/18.
 XX
 PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
 PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
 later myeloid differentiation activity.
 XX
 PS Disclosure; Page 32; 82pp; English.
 CC This protein sequence given comprises the entire amino acid sequence of
 CC human erythropoietin (EPO). EPO leads to the maturation of erythrocytes
 CC and is therefore designated as a late myeloid differentiation factor.
 CC (MDP). Within the scope of the invention hybrid molecules were produced
 CC which contain at least a portion of an early MDP and at least a portion
 CC of a late MDP covalently linked. The EPO sequence given is effective
 CC within the scope of the invention in full or in a truncated version.
 CC Amino acids 7-161 act as a late MDP when recombined with an early MDP eg.
 CC II-3. These compounds can be used to promote hematopoiesis in a patient.
 CC The bonding of the early and late factors allows a very high conc. of
 CC late MDP at the surface of a cell which the early MDP is bound. It also
 CC allows the early MDA to act more specifically to stimulate only the
 CC desired lineage, thus reducing undesirable effects. These compounds are
 CC useful for treating anaemias of various origins eg renal failure and
 CC AIDS. It is easier to produce and administer one recombinant molecule
 CC rather than two separate molecules
 XX
 SQ Sequence 166 AA;
 Query Match 100.0%; Score 846; DB 1; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2. 2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 OY 1 APPRLICDSVRVLLYRLEAKEAENTTGCAEHCSNLENITVDPDKVNFYAWKMEVGQAA 60
 Db 1 APPRLICDSVRVLLYRLEAKEAENTTGCAEHCSNLENITVDPDKVNFYAWKMEVGQAA 60
 OY 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 Db 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 OY 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 Db 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 OY 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 Db 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 OY 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 Db 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 RESULT 19
 ID AAR23593
 DE standard; protein; 166 AA.
 XX
 AC AAR23593;
 XX
 DT 20-OCT-1992 (first entry)
 DE Recombinant hematopoietic molecule portion 2.
 XX
 DE Erythropoietin; EPO: erythrocytes; IL-3; hematopoiesis.
 XX
 OS Homo sapiens.
 XX
 PN WO9206116-A.
 XX
 PR 28-SEP-1990; 90US-00589958.
 XX
 PA (ORTH) ORTHO PHARM CORP.
 XX
 PT Rosen JT;
 XX
 DR WPI; 1992-150819/18.
 XX
 PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
 PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
 later myeloid differentiation activity.
 XX
 PS Disclosure; Page 32; 82pp; English.
 CC This protein sequence given comprises the entire amino acid sequence of
 CC human erythropoietin (EPO). EPO leads to the maturation of erythrocytes
 CC and is therefore designated as a late myeloid differentiation factor.
 CC (MDP). Within the scope of the invention hybrid molecules were produced
 CC which contain at least a portion of an early MDP and at least a portion
 CC of a late MDP covalently linked. The EPO sequence given is effective
 CC within the scope of the invention in full or in a truncated version.
 CC Amino acids 7-161 act as a late MDP when recombined with an early MDP eg.
 CC II-3. These compounds can be used to promote hematopoiesis in a patient.
 CC The bonding of the early and late factors allows a very high conc. of
 CC late MDP at the surface of a cell which the early MDP is bound. It also
 CC allows the early MDA to act more specifically to stimulate only the
 CC desired lineage, thus reducing undesirable effects. These compounds are
 CC useful for treating anaemias of various origins eg renal failure and
 CC AIDS. It is easier to produce and administer one recombinant molecule
 CC rather than two separate molecules
 XX
 SQ Sequence 166 AA;
 Query Match 100.0%; Score 846; DB 2; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2. 2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 OY 1 APPRLICDSVRVLLYRLEAKEAENTTGCAEHCSNLENITVDPDKVNFYAWKMEVGQAA 60
 Db 1 APPRLICDSVRVLLYRLEAKEAENTTGCAEHCSNLENITVDPDKVNFYAWKMEVGQAA 60
 OY 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 Db 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 OY 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 Db 61 VEWVQGLALLSEAVLRGQALLVNSQWPWPLQHVDKAVSGLRSLTTLRAIGAQKEAIS 120
 OY 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 Db 121 PPDASAAPLRTTADTPRKLFVYSNPLRGKLYTGEACTGD 165
 RESULT 20
 ID AAW58404
 DE standard; protein; 166 AA.
 XX
 AC AAW58404;
 XX
 DT 12-OCT-1998 (first entry)
 DE Human erythropoietin.
 XX
 DE Erythropoietin receptor agonist; EPO; human; anaemia;
 KW hematopoietic deficiency; red blood cell; erythroid progenitor;
 KW bone marrow suppression.
 XX

OS Homo sapiens.
 XX KW Haematopoietic receptor agonist; erythropoietin receptor agonist; EPO;
 PN KW human; chimeric protein; stem cell expansion; tumour; infection;
 XX KW autoimmune disease; haematopoietic disorder; therapy; dendritic cell.
 PD 07-MAY-1998.
 XX
 PF 23-OCT-1997; 97WO-US018703.
 XX PR 25-OCT-1997; 96US-0034044P.
 PA (SEARL) SEARL & CO G D.
 XX
 PT McWherter CA, Feng Y, Summers N;
 XX DR WPI: 1998-272221/24.
 DR N-PSDB; AAV31031.
 PT Human erythropoietin receptor agonist polypeptide - used to stimulate the
 production of red blood cells in a patient.
 XX PS Claim 1; Page 93; 112pp; English.
 CC A claimed human erythropoietin (EPO) receptor agonist polypeptide
 comprises a modified EPO amino acid sequence given in AAW8404, where (a)
 optionally 1-6 amino acids from the N-terminus and 1-5 from the C-
 terminus can be deleted, (b) the N-terminus is joined to the C-terminus
 directly or through a linker (see AAW8405-12) capable of joining the N-
 terminus to the C-terminus, (c) there are new C- and N-termini at any two
 consecutive amino acids from amino acids 23-24 to 38-39, 40-41 to 41-42,
 43-44 to 48-49, 50-51 to 57-58, 77-78 to 82-83, 84-85 to 88-89, and 108-
 109 to 131-132, and (d) optionally the agonist polypeptide is preceded by
 Met, Ala, or Met-Ala. 60 Of these circularly permuted EPO receptor
 agonists (see AAW8413-72) are claimed. Also claimed are: nucleic acid
 molecules (see AAV30971-V31030) encoding novel EPO receptor agonists; a
 method of producing an EPO receptor agonist using transformed or
 transfected host cells; and methods for stimulating the production of
 haematopoietic cells, for selective ex vivo expansion of erythroid
 progenitors, and treating patients having a haematopoietic disorder using
 the EPO receptor agonists. The EPO receptor agonists retain one or more
 activities of native EPO and may also show improved haematopoietic cell-
 stimulating activity and/or an improved activity profile which may
 include reduction of undesirable biological activities associated with
 native EPO and/or have improved physical properties such as increased
 solubility, stability and refold efficiency
 XX SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 2; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2; e-86; Length 166;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRRLCDSRYLERTYLEAAGAENTTGCLEHCSLENNTVUPDTKVNFKRMEVGQA 60
 Db 1 APPRLCDSSYLERLILEAKRGAENTTGCLEHCSLENNTVUPDTKVNFKRMEVGQA 60
 QY 61 VENMGLALISEAVERGQALLVNSQWPQLRHDKAVSGRLSLTLLRGAQEKAIS 120
 Db 61 VENMGLALISEAVERGQALLVNSQWPQLRHDKAVSGRLSLTLLRGAQEKAIS 120
 QY 121 PPDAAASAPRITADTFRLFRVSNFLGKLUKTGEACRTG 165
 Db 121 PPDAAASAPRITADTFRLFRVSNFLGKLUKTGEACRTG 165

RESULT 21
 AAW7780 AAW7780 standard; protein; 166 AA.
 XX
 AAW7780; AC XX
 DT 24-NOV-1998 (first entry)
 DB Human EPO receptor agonist polypeptide.

XX
 KW Haematopoietic receptor agonist; erythropoietin receptor agonist; EPO;
 XX KW human; chimeric protein; stem cell expansion; tumour; infection;
 KW autoimmune disease; haematopoietic disorder; therapy; dendritic cell.
 OS Homo sapiens.
 XX
 FH Key location/Qualifiers
 FT Misc-difference 1. .6 /note= "possible positions of the N-terminus are optionally
 deleted"
 FT Misc-difference 23. .24 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 24. .25 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 25. .26 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 26. .27 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 27. .28 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 28. .29 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 29. .30 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 30. .31 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 31. .32 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 32. .33 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 33. .34 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 34. .35 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 35. .36 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 36. .37 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 37. .38 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 38. .39 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 39. .40 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 40. .41 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 41. .42 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 42. .43 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 43. .44 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 44. .45 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 45. .46 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 46. .47 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 47. .48 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 48. .49 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 49. .50 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 50. .51 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 51. .52 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 52. .53 /note= "possible positions of new C- and N-termini"
 FT Misc-difference 53. .54 /note= "possible positions of new C- and N-termini"

FT Misc-difference 54. .55
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 55. .56
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 56. .57
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 57. .58
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 77. .78
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 78. .79
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 79. .80
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 81. .82
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 82. .83
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 84. .85
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 85. .86
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 86. .87
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 87. .88
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 88. .89
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 108. .109
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 109. .110
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 110. .111
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 111. .112
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 112. .113
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 113. .114
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 114. .115
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 115. .116
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 116. .117
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 117. .118
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 118. .119
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 119. .120
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 120. .121
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 121. .122
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 122. .123
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 123. .124
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 124. .125
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 125. .126
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 126. .127
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 127. .128
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 128. .129
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 129. .130
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 130. .131
 FT /note= "possible positions of new C- and N-termini"

FT Misc-difference 131. .132
 FT /note= "possible positions of new C- and N-termini"
 FT Misc-difference 162. .166
 FT /note= "possible positions of new C- and N-termini"
 FT deleted" 1.5 amino acids of the C-terminus are optionally
 FT XX
 FT PN WQ981810-A2.
 FT XX 30-APR-1998.
 FT XX 23-OCT-1997; 97WO-US20037.
 FT XX PR 25-OCT-1996; 96US-0029629P.
 FT XX DR (SBAR) SEARLE & CO G D.
 FT XX PI McWherter CA, Feng Y, McKearn JP, Summers NL, Staten NR;
 FT XX Streeter PR, Minnerly JC, Minster NI, Woulfe SL;
 FT XX WPI; 1998-261504/23.
 FT PS Multi-functional chimeric haematopoietic receptor agonist - useful to
 FT treat haematopoietic disorders, tumours, infections or autoimmune
 FT diseases.
 FT XX Claim 1; Page 762; 841pp; English.
 CC A human erythropoietin (EPO) receptor agonist polypeptide comprises a
 CC modified EPO amino acid sequence of the formula provided in AAW7780, in
 CC which the N-terminus is joined to the C-terminus directly or via a
 CC linker, the polypeptide having new C- and N-termini at one of the
 CC positions indicated. Novel claimed multi-functional chimeric
 CC haemopoietic receptor agonists (see AAW7782-22) have the formula R1-L1
 CC -R2, R1-L1-R1, R1-R2 or R2-R1, where L is a linker and R1 and R2 are
 CC independently selected from: (a) the human EPO receptor agonist; (b)
 CC human stem cell factor receptor agonist polypeptide (see AAW7781); (c) a
 CC human fit-3 receptor agonist polypeptide (see AAW7782); (d) a modified
 CC human granulocyte colony stimulating factor (G-CSF) polypeptide (see
 CC AAW7783); (e) modified human interleukin-3 polypeptide (see AAW7784);
 CC (f) modified human c-mpl ligand polypeptide (see AAW7785); and (g) a
 CC factor selected from the group consisting of a CSF, a cytokine, a
 CC lymphokine, an interleukin and a haematopoietic growth factor, provided
 CC that at least R1 or R2 is selected from (a), (b) or (c) as above. The
 CC multi-functional chimeric haematopoietic receptor agonist can be used to
 CC stimulate the production of haematopoietic cells in a patient, for the ex
 CC vivo expansion of haematopoietic cells, for the production of dendritic
 CC Query Match 100%; Score 846; DB 2; Length 166;
 CC Best Local Similarity 100.0%; Pred. No. 2.2e-6; Indele 0; Gaps 0;
 CC Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;
 QY 1 APPRLCDSRVLLRVLRAKEAENITTGCAHCSLNENITPDTKNPYAWRMEVQQA 60
 DB 1 APPRLCDSRVLLRVLRAKEAENITTGCAHCSLNENITPDTKNPYAWRMEVQQA 60
 QY 61 VEWNGQALLSLRVLRAKEAENITTGCAHCSLNENITPDTKNPYAWRMEVQQA 120
 DB 61 VEWNGQALLSLRVLRAKEAENITTGCAHCSLNENITPDTKNPYAWRMEVQQA 120
 QY 121 PDDAASAPLRTTADPFRKLFVWSNPLRGKLYTGEACRTGD 165
 DB 121 PDDAASAPLRTTADPFRKLFVWSNPLRGKLYTGEACRTGD 165
 RESULT 22
 ABB07030
 ID ABB07030 standard; protein; 166 AA.
 XX
 ABB07030;
 AC
 XX
 DT 21-JUN-2002 (first entry)
 XX

XX
 PS Example 1; Page 22; 58pp; English.
 XX
 CC The present sequence is human erythropoietin (EPO) mature protein. EPO
 CC has improved biological activity and an extended serum half life greater
 CC than 20 hours. The present invention relates to modified EPOs such
 as fusion proteins comprising a Fc portion of an immunoglobulin (Ig)
 molecule and an EPO molecule (Fc-EPO). The Fc portion is fused covalently
 through its C-terminus directly or indirectly to the EPO molecule, and
 where the Fc portion as well as EPO portion may be modified or mutated.
 CC
 CC pattern of cysteines or disulphide bonding which is distinct from human
 CC or animal EPO. Pharmaceutical compositions containing EPO are useful in
 CC the treatment of EPO deficient diseases such as anaemia, renal failure,
 CC HIV infection, blood loss and chronic disease that can be treated with
 CC haematopoietic growth factor
 XX
 Sequence 166 AA;

Query Match 100.0%; Score 846; DB 4; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRYLRAKARENITGCAECSLNENITVDPDKYNFYAWKRMVGQAA 60
 Db 1 APPRLICDSRVLYLLEAKRAENITGCAECSLNENITVDPDKYNFYAWKRMVGQAA 60
 QY 61 VEWVOGLALLSBAVLKGQLVNSQPWEPLQHVDKAVSGRLSLTLLRAGQKAIS 120
 Db 61 VEWVOGLALLSBAVLKGQLVNSQPWEPLQHVDKAVSGRLSLTLLRAGQKAIS 120
 QY 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 QY 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165

RESULT 25
 AAB6698
 ID AAB6698 standard; protein; 166 AA.
 XX
 AC AAB6698;
 XX
 DT 06-APR-2001 (first entry)

DE Human erythropoletin protein #2.

KW Erythropoietin; EPO; reticulocytes; red blood cell; ethylene glycol;
 KW chronic renal failure; AIDS; cancer.
 OS Homo sapiens.
 XX
 WO200102017-A2.
 XX
 DD 11-JAN-2001.
 XX
 PP 28-JUN-2000; 2000WO-EP006009.
 XX
 PR 02-JUL-1999; 99US-0142243P.
 PR 05-AUG-1999; 99US-0147452P.
 PR 30-AUG-1999; 99US-0151454P.
 (HOFF) HOFFMANN LA ROCHE & CO AG F.
 PT Burg J, Hilger B, Josef H;
 XX
 DR WPI; 2001-147051/15.

XX
 PT Novel erythropoietin-glycoprotein conjugate useful for treating diseases
 PT correlated with anemia in chronic renal failure patients, AIDS and for
 PT treating cancer, is linked to polyethylene glycol through linker.
 XX
 PS Claim 19; Fig 2; 40pp; English.

CC The present invention relates to a conjugate comprising, human
 CC erythropoietin glycoprotein (EPO) having at least one free amino group
 CC and having in vivo biological activity of causing an increase the
 CC production of reticulocytes and red blood cells, covalently linked to 1-3
 CC lower-alkoxy poly(ethylene glycol) groups through a linker. The invention
 CC is useful for preparation of medicamenta for the treatment of prophylaxis
 CC of disease correlated with anemia in chronic renal failure patient^b
 CC (CRF), AIDS and for the treatment of cancer patients undergoing
 CC chemotherapy
 XX
 Sequence 166 AA;

Query Match 100.0%; Score 846; DB 4; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLYLLEAKRAENITGCAECSLNENITVDPDKYNFYAWKRMVGQAA 60
 Db 1 APPRLICDSRVLYLLEAKRAENITGCAECSLNENITVDPDKYNFYAWKRMVGQAA 60
 QY 61 VEWVOGLALLSBAVLKGQLVNSQPWEPLQHVDKAVSGRLSLTLLRAGQKAIS 120
 Db 61 VEWVOGLALLSBAVLKGQLVNSQPWEPLQHVDKAVSGRLSLTLLRAGQKAIS 120
 QY 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 QY 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFVYSNPLRGKLKLYGEACRTGD 165

RESULT 26
 ARBG92101
 ID ARBG92101 standard; protein; 166 AA.
 XX
 AC ARBG92101;
 XX
 DT 29-NOV-2002 (first entry)

XX
 DB Human erythropoietin (EPO).
 XX
 KW Human; erythropoietin; EPO; immunogenic; MHC class I; T-cell;
 KW major histocompatibility complex.
 OS Homo sapiens.
 XX
 PN WO200262843-A2.
 XX
 PD 15-AUG-2002.
 PP 05-FEB-2002; 2002WO-EP001174.
 XX
 PR 06-FEB-2001; 2001EP-00102615.
 PR 19-FEB-2001; 2001EP-00103954.
 XX
 PA (MERB) MERCK PATENT GMBH.
 XX
 Carr RJ, Carter G, Jones T, Williams S;
 XX
 DR WPI; 2002-62753/67.

PT New modified molecule that is non-immunogenic and which has the
 PT biological activity of human erythropoietin, useful for reducing
 PT propensity of the polypeptide to elicit an immune response upon
 PT administration to human subject.
 XX
 PS Disclosure; Page 5; 33pp; English.

XX
 CC The invention relates to a modified molecule having the biological
 CC activity of human erythropoietin (EPO) and being substantially non-
 CC immunogenic or less immunogenic than any non-modified molecule having the
 CC same biological activity when used in vivo. The modified molecule is
 CC useful for reducing propensity of the polypeptide to elicit an immune
 CC response upon administration to human subject. The 13mer T-cell group
 CC peptides having a potential MHC class II binding activity and created

CC from immunogenically non-modified erythropoietin, are useful for the
 CC manufacture of erythropoietin having substantially no or less
 CC immunogenicity than any non-modified molecule with the same biological
 CC activity when used in vivo. ABG92101-ABG92172 represent human
 CC erythropoietin and erythropoietin T-cell group peptides of the invention
 XX SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 5; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 APPRLIDSRSVRLERYLLEAKEAEKANITTCGAECISLNENITVPTDKTNYAWKRMEMVGQQA 60
 Db 1 APPRLIDSRSVRLERYLLEAKEAEKANITTCGAECISLNENITVPTDKTNYAWKRMEMVGQQA 60
 QY 61 VEWQGLALLSEAVLRLGQALLVNSQWEPPLQLHVDKAVSGRSLSLTIRALGAQEKAIS 120
 Db 61 VEWQGLALLSEAVLRLGQALLVNSQWEPPLQLHVDKAVSGRSLSLTIRALGAQEKAIS 120
 QY 121 PPDAAASAPLRTTADTFRKLRVYSNPLRGKLYTGEACTGD 165
 Db 121 PPDAAASAPLRTTADTFRKLRVYSNPLRGKLYTGEACTGD 165

RESULT 27

AM53062 ID AAM53062 standard; protein; 166 AA.

XX AC AAM53062;

XX DT 25-MAR-2002 (first entry)

DB Human erythropoietin (hEPO), 166 residue form.

XX Human; erythropoietin; EPO; hEPO; haemostatic; red blood cell;
 KW blood disorder; anaemia; chronic renal failure; CRP; AIDS;
 KW acquired immunodeficiency syndrome; cancer chemotherapy; haemostatic;
 KW anti-HIV; antianaemic.

OS Homo sapiens.

XX FH Key Location/Qualifiers

XX FT Disulfide-bond 7..161

FT Modified-site /note= "N-glycosylated"

FT Disulfide-bond 29..33

FT Modified-site 38

FT Modified-site 83

FT Modified-site 126

FT /note= "O-glycosylated"

XX SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 5; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLIDSRSVRLERYLLEAKEAEKANITTCGAECISLNENITVPTDKTNYAWKRMEMVGQQA 60
 Db 1 APPRLIDSRSVRLERYLLEAKEAEKANITTCGAECISLNENITVPTDKTNYAWKRMEMVGQQA 60
 QY 61 VEWQGLALLSEAVLRLGQALLVNSQWEPPLQLHVDKAVSGRSLSLTIRALGAQEKAIS 120
 Db 61 VEWQGLALLSEAVLRLGQALLVNSQWEPPLQLHVDKAVSGRSLSLTIRALGAQEKAIS 120
 QY 121 PPDAAASAPLRTTADTFRKLRVYSNPLRGKLYTGEACTGD 165
 Db 121 PPDAAASAPLRTTADTFRKLRVYSNPLRGKLYTGEACTGD 165

XX RESULT 28

ID ABB77897 standard; protein; 166 AA.

XX AC ABB77897;

XX DT 07-OCT-2002 (first entry)

XX DE Amino acid sequence of a human erythropoietin (EPO).

XX Human; erythropoietin; EPO; glycoprotein; reticulocyte production;
 KW red blood cell production; anaemia; chronic renal failure;
 KW acquired immunodeficiency syndrome; AIDS; cancer; bone marrow;
 KW committed erythroid progenitor.

OS Homo sapiens.

XX PN WO200187329-A1.

XX PD 22-NOV-2001.

XX PR 08-MAY-2001; 2001WO-EP005187.

XX PR 15-MAY-2000; 2000EP-00110355.

XX PR (HOFFMANN LA ROCHE & CO AG F.

XX PI Papadimitriou A;

XX DR WPI; 2002-082943/11.

XX PT Composition useful in the treatment of e.g. AIDS comprises an
 PT erythropoietin protein, and a multiple charged inorganic anion in a
 buffer.

XX PS Claim 28; Fig 2; 64pp; English.

PI Burg J, Engel A, Franz R, Hilger B, Schuring HB, Tischer W;
 PI Wozny M;
 XX
 WPI; 2002-566640/60.

PT Novel conjugate of erythropoietin glycoprotein with polyethylene glycol,
 useful for treating disease correlated with anemia in chronic renal
 failure patients and acquired immunodeficiency syndrome.

PS Claim 26; Fig 2; 40pp; English.

CC The present sequence represents a human erythropoietin (EPO) protein. It
 was used to produce conjugates of the invention. The specification
 describes a conjugate comprising an EPO glycoprotein having an N-terminal
 alpha-amino group, chosen from human EPO (hEPO) or its analogue (where
 hEPO is modified by addition of 1-6 glycosylation sites or a
 rearrangement of a glycosylation site). The glycoprotein is covalently
 linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo
 biological activity of causing bone marrow cells to increase production
 of reticulocytes and red blood cells. The conjugate increased circulating
 half-life and plasma residence time, decreased clearance, increased
 clinical activity in vivo, improved potency and stability, when compared
 to unmodified EPO. The EPO conjugate is useful for preparing medicaments
 for the treatment and prophylaxis of diseases correlated with anemia in
 chronic renal failure patients (CRF), acquired immunodeficiency syndrome
 (AIDS) and for treating cancer patients undergoing chemotherapy. It is
 also useful for treating patients by stimulating the division and
 differentiation of committed erythroid progenitors in the bone marrow
 sequence 166 AA;

Query Match 100%; Score 846; DB 5; Length 166;

Best Local Similarity 100.0%; Pred. No. 2_2e-66; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 APRLICDSRVIERYLLEAKEAENITGCAEHCISLNENITVDPDKVNIFYAWRRMEVGQQA 60
 Db 1 APRLICDSRVIERYLLEAKEAENITGCAEHCISLNENITVDPDKVNIFYAWRRMEVGQQA 60
 Oy 61 VEWQGLALLSEAVLRGQALLNSSSQEPHLQHDKAVSGRSLSLTIRALGAQEAIS 120
 Db 61 VEWQGLALLSEAVLRGQALLNSSSQEPHLQHDKAVSGRSLSLTIRALGAQEAIS 120
 Oy 121 PPDAAASAPLRTTADTPRKLFRVVSNFLRGKLUKYTGACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFRVVSNFLRGKLUKYTGACRTGD 165

RESULT 29

ID ADG65661
 ID ADG65661 standard; protein; 166 AA.

AC ADG65661;
 AC
 DT 11-MAR-2004 (first entry)

DE XX
 DE Human erythropoietin.

XX human; mouse; T-cell epitope; major histocompatibility complex; MHC;
 KW immunogenicity; MHC class II; antibody.
 Homo sapiens.

XX WO200269232-A2.

PD 06-SEP-2002.
 XX
 PR 18-FEB-2002; 2002WO-EP001688.

XX
 PR 19-FEB-2001; 2001EP-00103954.
 PR 08-MAR-2001; 2001EP-00105777.
 PR 15-MAR-2001; 2001EP-00106536.

PR 15-MAR-2001; 2001EP-00106538.

PR 20-MAR-2001; 2001EP-00106899.
 PR 20-MAR-2001; 2001EP-00107012.
 PR 27-MAR-2001; 2001EP-00107568.
 PR 25-APR-2001; 2001EP-00107220.
 PR 30-MAY-2001; 2001EP-00113228.
 PR 19-OCT-2001; 2001EP-00124955.
 PR 12-NOV-2001; 2001EP-00126859.
 XX
 PA (MERB) MERCK PATENT GMBH.

XX
 PI Carr PJ, Carter G, Jones T, Williams S, Hamilton A;
 DR XX
 DR WPI; 2002-750424/81.

PS Identifying potential T-cell epitope peptides within the amino acid
 sequence of a biological molecule, useful for preparing a biological
 molecule with reduced immunogenicity, comprises determining peptide
 PT binding to MHC molecules.

XX
 CC Example 7; Page 36; 85pp; English.

CC The invention relates to a novel method for identifying one or more
 potential T-cell epitope peptides within the amino acid sequence of a
 biological molecule by determining the binding of the peptides to major
 histocompatibility complex (MHC) molecules using in vitro or in silico
 techniques or biological assays. The method of the invention is useful
 for preparing a polypeptide, a protein, a fusion protein, an antibody or
 their fragments with reduced immunogenicity. The potential T-cell epitope
 peptide within the amino acid sequence of a parent immunogenically non-
 modified biological molecule identified is useful for preparing a
 biological molecule with reduced immunogenicity and having a retained
 desired biological activity, where the T-cell epitope is a 13mer peptide.

CC The present sequence is used in the exemplification of the invention.

XX Sequence 166 AA;

Query Match 100%; Score 846; DB 5; Length 166;
 Best Local Similarity 100%; Pred. No. 2_2e-66; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 APRLICDSRVIERYLLEAKEAENITGCAEHCISLNENITVDPDKVNIFYAWRRMEVGQQA 60
 Db 1 APRLICDSRVIERYLLEAKEAENITGCAEHCISLNENITVDPDKVNIFYAWRRMEVGQQA 60
 Oy 61 VEWQGLALLSEAVLRGQALLNSSSQEPHLQHDKAVSGRSLSLTIRALGAQEAIS 120
 Db 61 VEWQGLALLSEAVLRGQALLNSSSQEPHLQHDKAVSGRSLSLTIRALGAQEAIS 120
 Oy 121 PPDAAASAPLRTTADTPRKLFRVVSNFLRGKLUKYTGACRTGD 165
 Db 121 PPDAAASAPLRTTADTPRKLFRVVSNFLRGKLUKYTGACRTGD 165

RESULT 30

ABR3996
 ID ABR3996 standard; protein; 166 AA.
 XX
 AC ABR3996;
 XX
 DT 02-SEP-2003 (first entry)

XX Human erythropoietin (EPO) sequence.
 XX EPO; erythropoietin; mutuin; reticulocyte; red blood cell; antianemic;
 KW AIDS; cancer.
 XX
 OS Homo sapiens.

XX
 Key Location/Qualifiers
 FT Disulfide-bond 7..161
 /note= "disulphide bridge"
 FT Disulfide-bond 29..33
 /note= "disulphide bridge"

PT Modified-site 38
 FT /note= "Asn is N-glycosylated"
 FT Modified-site 83
 FT /note= "Asn is N-glycosylated"
 FT Modified-site 126
 FT /note= "Ser is O-glycosylated"
 XX PN WO2003029291-A2.
 XX PD 10-APR-2003.
 XX PF 20-SEP-2002; 2002WO-EP010556.
 XX PR 25-SEP-2001; 2001EP-00122555.
 XX PA (HOPP) HOFFMANN LA ROCHE & CO AG F.
 XX PT Tischer W;
 XX DR WPI; 2003-457226/43.
 XX PT Novel erythropoietin mutein having in vivo biological activity of causing bone marrow cells to increase production of reticulocytes/red blood cells, is N-glycosylated at Asn38 and Asn83 but not N-glycosylated at Asn24.
 XX PT Claim 6; Page 22; 22pp; English.
 PS XX The invention relates to an erythropoietin mutein (I) having the in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells, characterized by being N-glycosylated at Asn38 and Asn83 but not N-glycosylated at Asn24. (I) or an aqueous composition comprising an erythropoietin mutein is useful for the preparation of a medicament for the treatment or prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), AIDS and for the treatment of cancer patients undergoing chemotherapy. (I) or the composition is useful for treating a human patient experiencing blood disorders characterized by low or defective red blood cell production. (I) is useful for enhancing red blood cell formation. The present sequence represents a human erythropoietin (EPO) sequence.
 XX SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 6; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRILICDSVRLERYLLEAKEAENITGCAEHCSLENINITYPTKVNIFYAWKMEVGQAA 60
 Db 1 APPRICDSVRLERYLLEAKEAENITGCAEHCSLENINITYPTKVNIFYAWKMEVGQAA 60
 QY 61 VEWMOGLALLSEAVTRGQALLVNSQWPWPLQHVDKAVSGRLSITLRALGQKEAIS 120
 Db 61 VEWMOGLALLSEAVTRGQALLVNSQWPWPLQHVDKAVSGRLSITLRALGQKEAIS 120
 QY 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYTGEACRTG 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYTGEACRTG 165

RESULT 31
 ABR57500 ID ABR57500 standard; protein; 166 AA.
 AC ABR57500;
 XX DT 19-SEP-2003 (first entry)
 XX DE Human erythropoietin (EPO) amino acid sequence SEQ ID NO:1.
 XX KW Human; erythropoietin; EPO; hEPO; tranquilliser; cerebroprotective; anticonvulsant; vasotropic; antiinflammatory; immunosuppressive; antianaemic; antirheumatic; antiarthritic; anti-HIV; nephrotropic;
 KW KW red blood cell production stimulator; head trauma; stroke; epilepsy; ischaemia; hypoxia; immune-mediated inflammation; CNS disorder; HIV; excessive neuronal excitation; central nervous system disorder; chronic renal failure; anaemia; chronic inflammatory disease; rheumatoid arthritis.

XX OS Homo sapiens.
 XX PN WO200305526-A2.
 XX PD 10-JUL-2003.
 XX PR 18-DEC-2002; 2002WO-DR000871.
 XX PR 21-DEC-2001; 2001DK-00001953.
 XX PR 21-DEC-2001; 2001US-0343501P.
 XX PA (MAXY-) MAXXYGEN APS.
 XX PA (MAXY-) MAXXYGEN HOLDINGS LTD.
 XX PT Andersen KV;
 XX DR WPI; 2003-577388/54.
 XX PT Polypeptide conjugate useful in the treatment of e.g. stroke, head trauma and hypoxia comprises polymer molecule covalently attached to attachment site of human erythropoietin-like polypeptide.
 PS XX Disclosure; Page 61-62; 62pp; English.
 CC The present invention describes a polypeptide conjugate (I), which comprises at least one polymer molecule (a), covalently attached to an attachment site of a human erythropoietin-like polypeptide (b), where (b) comprises at least one removed and/or introduced lysine, cysteine, aspartic acid or glutamic acid residue compared to the amino acid sequence of human erythropoietin (hEPO). Also described: (1) a polypeptide comprising the amino acid sequence of (b); and (2) use of (1) as a pharmaceutical and in the preparation of a medicament for the prevention or treatment of disorders involving low or defective red blood cell production. (I) has tranquiliser, cerebroprotective, anticonvulsant, vasotropic, antiinflammatory, immunosuppressive, antianaemic, antirheumatic, antiarthritic, anti-HIV and nephrotropic activities, and can be used as a red blood cell production stimulator. (I) can be used as a pharmaceutical; in the manufacture of a medicament for prevention or treatment of disorders involving low or defective red blood cell production; and in the treatment of head trauma, stroke, epilepsy, ischaemia, hypoxia, immune-mediated inflammation, excessive neuronal excitation and other central nervous system (CNS) related conditions. Also useful for the treatment of HIV, chronic renal failure, anaemia in patients with non-myeloid malignancies, chronic inflammatory disease e.g. rheumatoid arthritis, anaemia associated with chronic disease, senile anaemia and anaemia in patients undergoing blood transfusion. The present sequence represents hEPO, which is given in the exemplification of the present invention
 XX SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 6; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2.2e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRILICDSVRLERYLLEAKEAENITGCAEHCSLENINITYPTKVNIFYAWKMEVGQAA 60
 Db 1 APPRICDSVRLERYLLEAKEAENITGCAEHCSLENINITYPTKVNIFYAWKMEVGQAA 60
 QY 61 VEWMOGLALLSEAVTRGQALLVNSQWPWPLQHVDKAVSGRLSITLRALGQKEAIS 120
 Db 61 VEWMOGLALLSEAVTRGQALLVNSQWPWPLQHVDKAVSGRLSITLRALGQKEAIS 120
 QY 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYTGEACRTG 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYTGEACRTG 165

QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DT	12-FEB-2004	(first entry)	
XX			
DB	Human erythropoietin (EPO).		
XX			
KW	immunostimulant; granulocyte macrophage colony stimulating factor; GM-CSF; neutropenia; myelosuppressive chemotherapy; bone marrow transplantation; HIV infection; burn; surgery; dilatation; anaemia; neonatal septicaemia; severe chronic neutropenia; aplastic anaemia; acute leukaemia; human; growth hormone super family; erythropoietin; EPO.		
KW	Homo sapiens.		
OS			
XX			
PN	US2003171284-A1.		
XX			
PD	11-SEP-2003.		
XX			
PF	15-NOV-2002; 2002US-0029814B.		
XX			
PR	14-JUL-1997; 97US-00522516P.		
PR	13-JUL-1998; 98WO-US014497.		
PR	14-JAN-2000; 2000US-00462941.		
PR	15-NOV-2001; 2001US-0332285P.		
PR	11-OCT-2002; 2002US-018040P.		
XX			
PA	(COXG/) COX G N.		
PA	(DOHR/) DOHERTY D H.		
XX			
PI	Cox GN, Doherty DH;		
XX			
DR	WPI; 2003-899295/82.		
XX			
PT	Protecting an animal from a disease or condition, useful for treating neutropenia, comprises administering to an animal having the disease or condition a composition comprising GM-CSF cysteine muttein.		
PT			
XX			
PS	Example 2; SEQ ID NO 2; 56pp; English.		
CC	The invention describes protecting an animal from a disease or condition that can be treated by wild-type granulocyte macrophage colony stimulating factor (GM-CSF) comprising administering to an animal having the disease or condition a composition comprising GM-CSF cysteine muttein.		
CC	The methods are useful for preventing or treating the occurrence of neutropenia in an animal, the neutropenia is selected from neutropenia resulting from myelosuppressive chemotherapy, neutropenia associated with bone marrow transplantation, neutropenia associated with infection with the human immunodeficiency virus, neutropenia associated with burns, surgery, dilatation, anaemia and neonatal septicaemia, severe chronic neutropenia, neutropenia associated with aplastic anaemia and acute leukaemia. This is the amino acid sequence of human erythropoietin (EPO), a member of the growth hormone super family which also includes interleukins.		
XX	Sequence 166 AA;		
SQ	Query Match 100.0%; Score 846; DB 7; Length 166; Best Local Similarity 100.0%; Prod. No. 2.2e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.		
Db	ADL92150 standard; protein; 166 AA.		
XX			
QY	121	PDDAASAAPLRTTADTRKLFRVYSNLRKGKULYGEACRTGD	165
DB	ADL92150 standard; protein; 166 AA.	</td	

Db	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD	165	DT	03-JUN-2004 (first entry)
RESULT 34			XX	Human cytokine protein #21.
ID ADK70564	ADK70564 standard; protein; 166 AA.		DE	Human cytokine; proteolysis; interferon; IFN; interleukin-10; IL-10;
ID ADK70564	ADK70564 standard; protein; 166 AA.		KW	Long-chain cytokine family; short-chain cytokine family; infection;
XX			KW	allergy; heart disease; cancer; liver disorder; autoimmune disease;
AC			KW	growth disorder; diabetes; neurodegenerative disease; antimicrobial;
XX			KW	antiallergic; cytostatic; immunosuppressive; antidiabetic;
DT	20-MAY-2004 (first entry)		KW	neuroprotective.
XX	Human erythropoietin (EPO) protein mature amino acid sequence.		OS	Homo sapiens.
XX	erythropoietin; EPO; non-immunogenic; immunogenic; EPO manufacture;		OS	Homo sapiens.
KW	erythropoietin; manufacture; anaemia; human.		OS	Homo sapiens.
XX			OS	Homo sapiens.
OS			OS	Homo sapiens.
PN	W02004018515-A2.		XX	W02004022593-A2.
XX			PN	
PD	04-MAR-2004.		XX	18-MAR-2004.
XX			PD	
PF	07-AUG-2003; 2003WO-EP008725.		XX	08-SEP-2003; 2003WO-1B004347.
XX			PR	09-SEP-2002; 2002US-040988P.
PR	09-AUG-2002; 2002EP-00017914.		PR	21-MAR-2003; 2003US-0457135P.
XX			XX	(NAUT-) NAUTILUS BIOTECH.
PA	(MERE) MERCK PATENT GMBH.		XX	
XX			PA	
PT	Baker M, Carr RJ;		XX	
XX			PI	Gantier R, Guyon T, Vega M, Drittanti L;
DR	DR		XX	
XX			DR	
PT	New modified human erythropoietin molecules with reduced immunogenicity,		XX	W01; 2004-248447/23.
PT	useful in various therapeutic applications such as in the treatment of		XX	
XX	anemia.		XX	
PS	Disclosure; Page 5; 38pp; English.		XX	
XX			XX	
CC	This invention relates to a novel modified molecule comprising the		CC	New modified cytokines with increased resistance to proteolysis, useful
CC	biological activity of human erythropoietin (EPO) and being substantially		CC	for diagnosing and treating diseases such as infections, allergies, heart
CC	non-immunogenic or less immunogenic than any non-modified molecule having		CC	diseases, cancer, liver disorders, autoimmune diseases or diabetes.
CC	the same biological activity in an individual when used in vivo. The		CC	Claim 88; SEQ ID NO 201; 316pp; English.
CC	invention is useful for manufacturing a modified human erythropoietin.		CC	The invention relates to modified cytokines that exhibit increased
CC	molecule. The modified EPO may be used in various therapeutic		CC	resistance to proteolysis compared to unmodified cytokines. The invention
CC	applications, such as in the treatment of anaemia. The present sequence		CC	also relates to nucleic acid molecules encoding the cytokines, a
CC	is that of the mature human erythropoietin protein which was used to		CC	pharmaceutical composition comprising a nucleic acid molecule in a
CC	derive the modified EPO molecules of the invention.		CC	pharmaceutical carrier, and a method of generating a protein or peptide
SQ	Sequence 166 AA;		CC	altered phenotype. The modified cytokine is selected from a member of the
Query Match	100.0%; Score 846; DB 8; Length 166;		CC	interferons (IFNs)/interleukin (IL)-10 protein family, a member of the
Best Local Similarity	100.0%; Pred. No. 2.2e-86;		CC	long-chain cytokine family or a member of the short-chain cytokine
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		CC	family. The composition and method are useful for diagnosing and treating
OY	1 APRPLICDSVRLEYLLEAKEAENITTGCAEHCSLNENITVDPDKVNFKVMEVGQAA 60		CC	diseases such as infections, allergies, heart diseases, cancer, liver
Db	1 APPRLICDSVRLEYLLEAKEAENITTGCAEHCSLNENITVDPDKVNFKVMEVGQAA 60		CC	disorders, autoimmune diseases, growth disorders, diabetes or
OY	61 VEWQGALLSLEAVIRGOALIINNSQPWBLQLHYDKAVSGRLSLTLLRALGAQKAI S 120		CC	neurodegenerative diseases. This sequence represents a human cytokine
Db	61 VEWQGALLSLEAVIRGOALIINNSQPWBLQLHYDKAVSGRLSLTLLRALGAQKAI S 120		CC	protein of the invention.
SQ	Sequence 166 AA;		XX	
Query Match	100.0%; Score 846; DB 8; Length 166;		XX	
Best Local Similarity	100.0%; Pred. No. 2.2e-86;		XX	
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		XX	
OY	1 APRPLICDSVRLEYLLEAKEAENITTGCAEHCSLNENITVDPDKVNFKVMEVGQAA 60		Db	1 APRPLICDSVRLEYLLEAKEAENITTGCAEHCSLNENITVDPDKVNFKVMEVGQAA 60
Db	1 APPRLICDSVRLEYLLEAKEAENITTGCAEHCSLNENITVDPDKVNFKVMEVGQAA 60		OY	61 VEWQGALLSLEAVIRGOALIINNSQPWBLQLHYDKAVSGRLSLTLLRALGAQKAI S 120
OY	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD 165		Db	61 VEWQGALLSLEAVIRGOALIINNSQPWBLQLHYDKAVSGRLSLTLLRALGAQKAI S 120
Db	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD 165		OY	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD 165
Db	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD 165		Db	121 PPDAAASAPRLTTADTRKLFRVSNFLRGKLUKYGEACRTGD 165
RESULT 35				
AD18867				
ID ADL8867	ADL8867 standard; protein; 166 AA.			
XX				
AC	ADL8867;			
XX				
RESULT 36				
AD106781				
ID ADL06781	ADL06781 standard; protein; 166 AA.			
XX				
AC	ADL06781;			
XX				

		ID	ADD59416 standard; protein; 166 AA.
XX		XX	
DT	03-JUN-2004 (first entry)	AC	ADD59416;
XX		XX	
DB	Human 166 residue erythropoietin (EPO), SEQ ID NO:2.	DT	26-AUG-2004 (first entry)
XX		XX	
KW	Human; erythropoietin; EPO; iron distribution disturbance; diabetes; non-insulin dependent diabetes; type 2 diabetes; reticulocyte production; red blood cell production; antidiabetic.	XX	Human 166 residue erythropoietin (EPO), SEQ ID NO:2.
KW		XX	
KW	Homo sapiens.	XX	Human; erythropoietin; EPO; iron distribution disturbance; heart disease; heart insufficiency; coronary heart disease; atherosclerosis; acute coronary syndrome; heart failure; congestive heart failure; reticulocyte production; red blood cell production; cardiologist; antiarrtherosclerotic.
OS		XX	
PN	WO2004019972-A1.	XX	
XX		XX	
XX	11-MAR-2004.	PD	10-JUN-2004.
XX		XX	
PP	20-AUG-2003; 2003WO-EP009194.	PR	17-NOV-2003; 2003WO-EP012822.
XX		XX	
XX	29-AUG-2002; 2002EP-00019100.	PR	22-NOV-2002; 2002EP-00026342.
XX		XX	
PA	(HOFF) HOFFMANN LA ROCHE & CO AG F.	PA	(HOFF) HOFFMANN LA ROCHE & CO AG F.
XX		XX	
PT	Lehmann P., Roeddiger R., Walter-Matsui R;	PT	Lehmann P., Roeddiger R., Walter-Matsui R;
XX		XX	
PS	DR; WPI; 2004-282643/26.	PS	DR; WPI; 2004-450212/42.
XX		XX	
CC	The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in diabetes. The	CC	The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in diabetes. The
CC	erythropoietin protein is preferably a human erythropoietin (e.g.,	CC	erythropoietin protein is preferably a human erythropoietin (e.g.,
CC	epoetin alpha and epoetin beta) which may be expressed by endogenous gene	CC	epoetin alpha and epoetin beta) which may be expressed by endogenous gene
CC	activation or an erythropoietin analogue such as darbepoietin alpha. The	CC	activation or an erythropoietin analogue such as darbepoietin alpha. The
CC	erythropoietin protein used in the method may also be modified by the	CC	erythropoietin protein used in the method may also be modified by the
CC	addition of 1-6 glycosylation sites, or by pegylation. Patients with	CC	addition of 1-6 glycosylation sites, or by pegylation. Patients with
CC	diabetes have been found to have a high probability of be affected by	CC	diabetes have been found to have a high probability of be affected by
CC	disturbances of iron distribution. In such patients, the overall	CC	disturbances of iron distribution. In such patients, the overall
CC	concentration of iron in the body is normal (compared with conditions	CC	concentration of iron in the body is normal (compared with conditions
CC	such as anaemia), but the individual may suffer the effects of iron	CC	such as anaemia), but the individual may suffer the effects of iron
CC	accumulation in certain organs, leading to organ damage and destruction,	CC	accumulation in certain organs, leading to organ damage and destruction,
CC	and/or experience effects similar to anaemia due to iron usage in blood	CC	and/or experience effects similar to anaemia due to iron usage in blood
CC	cell formation being impaired. Erythropoietin causes bone marrow cells to	CC	cell formation being impaired. Erythropoietin causes bone marrow cells to
CC	increase production of reticulocytes and red blood cells, and this has	CC	increase production of reticulocytes and red blood cells, and this has
CC	been found to have a beneficial effect on iron distribution disturbances	CC	been found to have a beneficial effect on iron distribution disturbances
CC	in diabetes e.g., non-insulin dependent (type 2) diabetes. Erythropoietin	CC	in diabetes e.g., non-insulin dependent (type 2) diabetes. Erythropoietin
CC	proteins may be used to manufacture a medicament for the	CC	proteins may be used to manufacture a medicament for the
CC	treatment of disturbances of iron distribution in diabetes. The present	CC	treatment of disturbances of iron distribution in diabetes. The present
CC	sequence represents a 166 amino acid human erythropoietin which is	CC	sequence represents a 166 amino acid human erythropoietin which is
CC	specifically claimed for use in the invention.	CC	specifically claimed for use in the invention.
XX		XX	
SQ	Sequence 166 AA;	SQ	Sequence 166 AA;
	Query Match 100.0%; Score 846; DB 8; Length 166;		Query Match 100.0%; Score 846; DB 8; Length 166;
	Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;		Best Local Similarity 100.0%; Pred. No. 2.2e-86; Mismatches 0; Indels 0; Gaps 0;
OY	1 APPRLICDSRVLRLVYLLLEAKEAENITTGCAEHCSLNENITVPTDKVNLYAWKRMEVGQQA 60	OY	1 APPRLICDSRVLRLVYLLLEAKEAENITTGCAEHCSLNENITVPTDKVNLYAWKRMEVGQQA 60
Db	1 APPRLICDSRVLRLVYLLLEAKEAENITTGCAEHCSLNENITVPTDKVNLYAWKRMEVGQQA 60	Db	1 APPRLICDSRVLRLVYLLLEAKEAENITTGCAEHCSLNENITVPTDKVNLYAWKRMEVGQQA 60
OY	61 VEWVQGLALLSEAVLVRQQLLVNSSQPWEPLQHVDAVGSLSLTILRALGAQEAI 120	OY	61 VEWVQGLALLSEAVLVRQQLLVNSSQPWEPLQHVDAVGSLSLTILRALGAQEAI 120
Db	61 VEWVQGLALLSEAVLVRQQLLVNSSQPWEPLQHVDAVGSLSLTILRALGAQEAI 120	Db	61 VEWVQGLALLSEAVLVRQQLLVNSSQPWEPLQHVDAVGSLSLTILRALGAQEAI 120
OY	121 PPDRASAAPLTTADTRKLFRYTSNPFNGKUKLYGRACRGD 165	OY	121 PPDRASAAPLTTADTRKLFRYTSNPFNGKUKLYGRACRGD 165
Db	121 PPDRASAAPLTTADTRKLFRYTSNPFNGKUKLYGRACRGD 165	Db	121 PPDRASAAPLTTADTRKLFRYTSNPFNGKUKLYGRACRGD 165
RESULT 37	ADO59416	QY	61 VEWVQGLALLSEAVLVRQQLLVNSSQPWEPLQHVDAVGSLSLTILRALGAQEAI 120

Qy 121 PPDAAASAPLTTADTFRKLFRVYNSFLRGKLUKYGEACRTGD 165
 Db 121 PPDAAASAPLTTADTFRKLFRVYNSFLRGKLUKYGEACRTGD 165

RESULT 38

ID ADV67303; standard; peptide; 166 AA.

XX

AC ADV67303;

XX

DT 10-MAR-2005 (first entry)

XX

DE Amino acid sequence of mature human erythropoietin.

XX

KW antianemic; antisickling; CNS-Gen; gynecological; neuroprotective; hematoologic irregularity; erythropoietin; EPO; EPO conjugate; anemia; cystic fibrosis; pregnancy; menstrual disorder; spinal cord injury.

OS Homo sapiens.

XX

PN WO2004108667-A2.

XX

PD 16-DEC-2004.

XX

PP 27-MAY-2004; 2004WO-US016670.

XX

PR 30-MAY-2003; 2003US-0475074P.

PA (CENZ) CENTOCOR INC.

XX

PT Pool CT;

XX

DR WPI; 2005-048518/05.

XX

PT Erythropoietin conjugate useful for treating anemia, has ability of causing bone marrow cells to increase production of red blood cells, and comprising moiety of erythropoietin, modified amino acids and organic moieties.

XX

PS Disclosure; SEQ ID NO 7; 41PP; English.

XX

The specification describes erythropoietin (EPO) conjugates, derived from formulas given in the specification (see Adv6729). These conjugates cause bone marrow cells to increase production of red blood cells. The EPO conjugates have increased serum half-life compared to unconjugated erythropoietin. EPO conjugates of the invention are useful for treating anemia, as well as a variety of disease states of hematologic irregularity e.g. sickle cell disease, beta-thalassemia, cystic fibrosis, pregnancy, menstrual disorder, and spinal cord injury. The present sequence represents mature human EPO.

XX

SQ Sequence 166 AA;

Query Match 100.0%; Score 846; DB 9; Length 166;
 Best Local Similarity 100.0%; Pred. No. 2-2e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDPRVLTLEAKAENITGCELEHCSLENNTVPDTKVNFTAWKRMVEGQDA 60
 1 APPRLICDPRVLTLEAKAENITGCELEHCSLENNTVPDTKVNFTAWKRMVEGQDA 60

Qy 61 VEWNGGLIISSEAVRQALLNSQWPBLQLHQDKAVSGLRSITTLRAGQEAIS 120
 61 VEWNGGLIISSEAVRQALLNSQWPBLQLHQDKAVSGLRSITTLRAGQEAIS 120

Db 121 PPDAAASAPLTTADTFRKLFRVYNSFLRGKLUKYGEACRTGD 165
 Qy 121 PPDAAASAPLTTADTFRKLFRVYNSFLRGKLUKYGEACRTGD 165
 Db 121 PPDAAASAPLTTADTFRKLFRVYNSFLRGKLUKYGEACRTGD 165

RESULT 39

ID ADY93798

XX

AC ADY93798;

XX

DT 02-JUN-2005 (first entry)

XX

DB Human erythropoietin protein SEQ ID NO:2.

XX

KW somatotropin; site-specific mutagenesis; antianemic; anemia.

OS Homo sapiens.

XX

PN US2005058621-A1.

XX

PD 17-MAR-2005.

XX

PP 13-OCT-2003; 2003US-0068588.

XX

PR 14-JAN-1997; 97US-0052516P.

PR 13-JUL-1998; 98WO-US014497.

PR 14-JAN-1999; 99US-011601P.

PR 14-JAN-2000; 2000US-00462241.

PR 14-JAN-2000; 2000WO-US000331.

PR 16-MAY-2000; 2000US-0204617P.

PR 16-MAY-2001; 2001WO-US016088.

PR 06-SEP-2001; 2001US-00889273.

PR 15-NOV-2001; 2001US-0332245P.

PR 11-OCT-2002; 2002US-0418040P.

PR 11-OCT-2002; 2002US-0418105P.

PR 11-OCT-2002; 2002US-041816P.

PR 15-NOV-2002; 2002US-0029848.

PR 26-MAR-2003; 2003US-00400377.

PR 10-APR-2003; 2003US-00276358.

XX

PA (COXG/) COX G N.

XX

PT Cox GN;

XX

DR WPI; 2005-312503/32.

XX

PT Protecting animal from disease or condition, e.g. neutropenia, anemia or cancer, that can be treated by granulocyte colony-stimulating factor, erythropoietin, or alpha interferon, comprises administering cysteine variant of the protein.

XX

PS Claim 18; SEQ ID NO 2; 66PP; English.

XX

The invention describes a method for protecting an animal from a disease or condition that can be treated by granulocyte colony-stimulating factor (G-CSF), erythropoietin (EPO) or alpha interferon-2. The method comprises administering to the animal a composition comprising a cysteine variant of G-CSF, EPO or alpha interferon. The method is useful for protecting an animal from a disease or condition that can be treated by G-CSF, where the disease is neutropenia. The neutropenia can be neutropenia resulting from chemotherapy, neutropenia associated with bone marrow transplantation, infection with HIV or burns, surgery, dilatation, anemia and neonatal septicemia, severe chronic neutropenia, and neutropenia associated with aplastic anemia and acute leukemia. The method is also useful for protecting an animal from a disease or condition that can be treated by EPO, where the disease is anemia. The anemia can be anemia resulting from chemotherapy, anemia resulting from renal disease, anemia resulting from renal failure and anemia resulting from drug complications. The method is also useful for protecting an animal from a disease or condition that can be treated by alpha interferon-2, where the disease is cancer or viral disease (preferably hepatitis B or hepatitis C). The present sequence represents the human EPO protein, which is used in an example from the present invention for creating cysteine-added variants of EPO.

SQ Sequence 166 AA;

Query	Match	100.0%;	Score	846;	DB	9;	Length	166;
Best Local Similarity	100.0%	;	Pred. No.	2.2e-86;				
Matches	165;	Conservative	0;	Mismatches	0;	Indels	0;	Gaps
OY								
Db	1	APRILICDSRVLYERVYLEAKERAENTITGCAEHCSLAMENTITPDTKWNVYAWKRMVGQOA	60					
	.1	APRILICDSRVLYERVYLEAKERAENTITGCAEHCSLAMENTITPDTKWNVYAWKRMVGQOA	60					
Qy	61	VEVMCGALLSEAVVRGQQLVNSSCPWPQLQHDKAVSGRLSLTTLAIGAQKEAIS	120					
Db	61	VEVMCGALLSEAVVRGQQLVNSSCPWPQLQHDKAVSGRLSLTTLAIGAQKEAIS	120					
Qy	121	PPDAASAPERTITADTFKLPRVSNFLAGKLUYTGEACRTGD	165					
Db	121	PPDAASAPERTITADTFKLPRVSNFLAGKLUYTGEACRTGD	165					
RESULT 40								
AEA47165	AEA47165 standard; protein; 166 AA.							
XX	XX							
AC	AC							
XX	XX							
DT	DT							
11-AUG-2005	(first entry)							
XX	XX							
DE	DE							
Brythropoietin peptide, SEQ ID No 2.								
XX	XX							
KW	erythropoietin; antianemic; vasoactive; antiinflammatory; blood; ischemia; trauma; inflammation.							
XX	XX							
OS	Synthetic.							
XX	XX							
FH	Key	Location/Qualifiers						
FT	Disulfide-bond	7. .161						
FT	Modified-site	24						
FT	Disulfide-bond	29. .33						
FT	Modified-site	38						
FT	Modified-site	83						
FT	Modified-site	126						
FT	Modified-site	/note= "Linked to CMP-Sialic acid-PEG structure"						
FT	Modified-site	/note= "N-glycosylated or O-phosphorylated"						
FT	Modified-site	/note= "N-glycosylated or O-phosphorylated"						
PN	WO2005051327-A2.							
XX	XX							
XX	XX							
XX	XX							
XX	XX							
PP	24-NOV-2004;	2004WO-US039712.						
PP	09-JUN-2005.							
DR								
PR	24-NOV-2003;	2003US-0524989P.						
PR	26-JAN-2004;	2004US-0539387P.						
PR	22-MAR-2004;	2004US-055504P.						
PR	23-TUL-2004;	2004US-0590573P.						
PR	29-JUL-2004;	2004US-0592744P.						
PR	29-SEP-2004;	2004US-0614518P.						
PR	29-OCT-2004;	2004US-0623387P.						
PA	(NEOS-)	NEOSE TECHNOLOGIES INC.						
PI	Defrees S,	Bayer RJ,	Zopf DA;					
XX	XX							
XX	XX							
PS	New Polyethyleneglycolated amino saccharide erythropoietin peptide useful for treating tissue injury e.g. damage resulting from ischemia, trauma, inflammation or contact with toxic substances.							
XX	XX							
PS	Disclosure; SBQ ID NO 2; 128pp; English.							
XX	XX							
CC	The invention relates to a novel polyethyleneglycolated amino saccharide erythropoietin peptide. The invention further comprises a method for making a polyethyleneglycolated amino saccharide erythropoietin peptide							

containing an amino sugar moiety of formula X-O-Gal-. The method involves contacting a substrate erythropoietin peptide comprising the Gal moiety with a polyethylene glycol (PEG)-sialic acid donor moiety of the above formula and an enzyme that transfers the PEG-sialic acid onto the Gal of the glycosyl moiety for the transfer. The novel polyethylene glycolated amino saccharide erythropoietin peptide has antianemic, vasotrophic, and antiinflammatory activities. The polyethylene glycolated amino saccharide erythropoietin peptide is useful for treating a condition, such as compromised red blood cell production and also tissue injury, such as damage resulting from ischemia, trauma, inflammation or contact with toxic substances in a subject. The modified erythropoietin peptide enhances red blood cell production, and can be prepared by a cost effective method. This sequence represents an exemplary erythropoietin peptide for use in the novel glycopegylated erythropoietin peptide of the invention.

RESULT 41
ABB21318
ID ABB21318 standard; protein; 166 AA.
XX
AC ABB21318;
XX
DT 08-SEP-2005 (first entry)
XX
DB Amino acid sequence of human erythropoietin #2.
XX
KW erythropoietin; pharmaceutical; gastrointestinal inflammation;
KW antiinflammatory; gastrointestinal-gen.; colitis; antiinflammatory;
KW gastrointestinal disease; inflammation.
XX
OS Homo sapiens.
XX
PN WO2005058347-A1.
XX
PD 30-JUN-2005.
PP 10-DEC-2004; 2004WO-EP014105.
XX
PR 19-DEC-2003; 2003EP-00104832.
XX
DA (HOFER) HOPFMANN LA ROCHE & CO AG F.
XX
PT Klma H, Lehmann P, Roeddiger R, Walter-Matsui R;
XX
DR
XX
PT Use of erythropoietin protein in manufacture of medicament for treatment
PT of disturbances of iron distribution in chronic inflammatory intestinal
PT diseases.
XX
PS Claim 7; SEQ ID NO 2; 32pp; English.
XX
CC The specification describes the use of erythropoietin protein in the
manufacture of medicament for the treatment of disturbances of iron

Db 62 VEWQGLALLSEAVLQRGQALLVNSSQWEPOLQHVDKAVSGLSLTLLRAGQKAIS 121
 Qy 121 PPDAAASAPLRTITADPFLKLFVRSVPLRGKLUYGEACRTGD 165
 -Best Local Similarity 100.0%; Pred. No. 2. 3-86;
 Db 122 PPDAAASAPLRTITADPFLKLFVRSVPLRGKLUYGEACRTGD 166
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 Result 44
 ABB77899
 ID ABB77899 standard; protein; 169 AA.
 XX
 AC ABB77899;
 XX
 DT 07-OCT-2002 (first entry)
 DE Amino acid sequence of a modified human erythropoietin (EPO).
 XX
 KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production;
 KW red blood cell production; anaemia; chronic renal failure;
 KW acquired immunodeficiency syndrome; AIDS; cancer; bone marrow;
 KW committed erythroid progenitor.
 XX
 OS Synthetic.
 XX
 Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Cleavage-site 1..3
 FT /note= "proteolytic cleavage site"
 FT Protein 4..174
 FT /note= "EPO protein"
 XX
 WO200249673-A2.
 XX
 PD 27-JUN-2002.
 XX
 PP 08-DEC-2001; 2001WO-EP014434.
 PI Burg J, Engel A, Franze R, Hilger B, Schurig HE, Tischer W;
 PI Wozny M;
 XX
 PR 20-DEC-2000; 2000EP-00127891.
 XX
 PA (HOFF) HOFFMANN LA ROCHE & CO AG F.
 XX
 DR 08-DEC-2001; 2001WO-EP014434.
 PR 20-DEC-2000; 2000EP-00127891.
 PS Disclosure; Page 39; 40PP; English.
 XX
 CC The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased circulating half-life and plasma residence time, decreased clearance, increased clinical activity in vivo, improved potency and stability, when compared to unmodified EPO. The EPO conjugate is useful for preparing medicaments for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRE), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow.
 CC Sequence 169 AA;

Db 61 VEWQGLALLSEAVLQRGQALLVNSSQWEPOLQHVDKAVSGLSLTLLRAGQKAIS 120
 Qy 64 VEWQGLALLSEAVLQRGQALLVNSSQWEPOLQHVDKAVSGLSLTLLRAGQKAIS 123
 DB 64 VEWQGLALLSEAVLQRGQALLVNSSQWEPOLQHVDKAVSGLSLTLLRAGQKAIS 123
 ABB77898
 ID ABB77898 standard; protein; 174 AA.
 XX
 AC ABB77898;
 XX
 DT 07-OCT-2002 (first entry)
 DE Amino acid sequence of a modified human erythropoietin (EPO).
 XX
 KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production;
 KW red blood cell production; anaemia; chronic renal failure;
 KW acquired immunodeficiency syndrome; AIDS; cancer; bone marrow;
 KW committed erythroid progenitor.
 XX
 OS Synthetic.
 XX
 Homo sapiens.
 XX
 FH Key Location/Qualifiers
 FT Cleavage-site 1..8
 FT /note= "proteolytic cleavage site"
 FT Protein 9..174
 FT /note= "EPO protein"
 XX
 WO200249673-A2.
 XX
 PR 20-DEC-2001; 2001WO-EP014434.
 PR 20-DEC-2000; 2000EP-00127891.
 PA (HOFF) HOFFMANN LA ROCHE & CO AG F.
 XX
 DR 08-DEC-2001; 2001WO-EP014434.
 PR 20-DEC-2000; 2000EP-00127891.
 PS Disclosure; Page 38-39; 40PP; English.
 XX
 CC The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased

CC circulating half-life and plasma residence time, decreased clearance, increased clinical activity in vivo, improved potency and stability, when compared to unmodified EPO. The EPO conjugate is useful for preparing medicaments for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

SQ Sequence 174 AA;

Query Match 100.0%; Score 846; DB 5; Length 174;
Best Local Similarity 100.0%; Pred. No. 2. 4e-86; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAEINTTGCACHECSLNENITVPTKVNLYAKWMEVGQA 60
Db 9 APPRLICDSRVLERYLLEAKEAEINTTGCACHECSLNENITVPTKVNLYAKWMEVGQA 60
QY 61 VEWMOGLALLSEAVLRGQALVNSSQWEPFLQHDKAVSGLSLTILRALGAQEAKIS 120
Db 69 VEWMOGLALLSEAVLRGQALVNSSQWEPFLQHDKAVSGLSLTILRALGAQEAKIS 120
QY 121 PPDASAASAPLRTTADTRKLFRVYSNPLRGKLYGEACTGD 165
Db 129 PPDASAASAPLRTTADTRKLFRVYSNPLRGKLYGEACTGD 173

RESULT 46

ABB77900

ID ABB77900 standard; protein; 174 AA.

XX ABB77900;

AC ABB77900;

XX DT 07-OCT-2002 (first entry)

XX DE Amino acid sequence of a modified human erythropoietin (EPO).

XX KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production; red blood cell production; anaemia; chronic renal failure; acquired immunodeficiency syndrome; AIDS; cancer; bone marrow; committed erythroid progenitor.

KW

OS Synthetic.

OS Homo sapiens.

XX FH Key Cleavage-site

FT 1..8 /note= "proteolytic cleavage site"

FT 9..174 /note= "EPO protein"

XX PN WO200249673-A2.

XX DD 27-JUN-2002.

PP XX 08-DEC-2001; 2001WO-EPO14434.

PP XX PR 20-DEC-2000; 2000EP-00127891.

PA (HOFFMANN LA ROCHE & CO AG F.

XX PA (HOFFMANN LA ROCHE & CO AG F.

PI Burg J, Engel A, Franze R, Hilger B, Schurig HE, Tischer W;

PI Wozny M;

XX DR WPI; 2002-566640/60.

XX PT Novel conjugate of erythropoietin glycoprotein with polyethylene glycol, useful for treating diseases correlated with anaemia in chronic renal failure patients and acquired immunodeficiency syndrome.

PT

PS Disclosure; Page 39-40; 40PP; English.

XX

The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased circulating half-life and plasma residence time, decreased clearance, compared to unmodified EPO. The EPO conjugate is useful for preparing with medicaments for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

SQ Sequence 174 AA;

Query Match 100.0%; Score 846; DB 5; Length 174;
Best Local Similarity 100.0%; Pred. No. 2. 4e-86; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAEINTTGCACHECSLNENITVPTKVNLYAKWMEVGQA 60
Db 9 APPRLICDSRVLERYLLEAKEAEINTTGCACHECSLNENITVPTKVNLYAKWMEVGQA 60
QY 61 VEWMOGLALLSEAVLRGQALVNSSQWEPFLQHDKAVSGLSLTILRALGAQEAKIS 120
Db 69 VEWMOGLALLSEAVLRGQALVNSSQWEPFLQHDKAVSGLSLTILRALGAQEAKIS 120
QY 121 PPDASAASAPLRTTADTRKLFRVYSNPLRGKLYGEACTGD 165
Db 129 PPDASAASAPLRTTADTRKLFRVYSNPLRGKLYGEACTGD 173

RESULT 47

AAP60599

ID AAP60599 standard; protein; 188 AA.

XX AAC AAP60599;

XX DT 25-MAR-2003 (revised)

DT 01-JAN-1980 (first entry)

XX DB C1 clone lambda HEPOL16 encoding human erythropoietin.

XX KW Erythropoietin; lambda HEPOL16; recombinant plasmid vector; anaemia; mammal cell culture; 3T3; CHO; Chinese hamster ovary; ss.

XX OS Homo sapiens.

XX PN WO860320-A.

XX DD 19-JUN-1986.

PP 03-DEC-1985; 85WO-US002405.

XX PR 04-DEC-1984; 84US-0057713.

PR 03-JAN-1985; 85US-0068822.

PR 22-JAN-1985; 85US-00693258.

XX PA (GEMY) GENETICS INST INC.

PA (FRIT/) FRITSCHIE E.

PI Fritsch E, Hewick RM, Jacobs K;

XX DR WPI; 1986-169459/26.

DR N-PSDB; AAN60519.

XX Prod'n. of human cDNA clone expressing erythropoietin - for mass prodn. of
 PT erythropoietin, useful for treating anaemia.
 XX Disclosure; Page 20; 61pp; English.

PS A recombinant plasmid vector expressing this clone is expressed in e. g.
 CC 3T3 or CHO cell cultures. The produced erythropoietin is useful for
 CC treatment of anaemia, especially renal anaemia. The cloned gene expresses
 CC high levels of the protein and thus provides a means of mass production.
 CC See also AAN60513-18, AAN60520-21 and AAF60598. (Updated on 25-MAR-2003
 CC to correct PA field.)
 XX

SQ Sequence 188 AA;

	Query Match	Score	DB 1;	Length	188;
Best Local Similarity	100.0%	Score	846;	DB 1;	Length 188;
Matches	165;	Pred. No.	2.7e-86;	Mismatches	0;
		Indels	0;	Gaps	0;

QY 1 APPRLCDSRVLLERYLLEAKAENITTCGAECNSLNENITVPDTKVNPFYAWKRMVEGQQA 60
 DB 23 APPRLCDSRVLLERYLLEAKAENITTCGAECNSLNENITVPDTKVNPFYAWKRMVEGQQA 82

QY 61 VEWMOGLALLSEAVLGRQALLVNSQWPFLQHVDKAVSGSLSLTTLRAGAQEAIS 120
 DB 83 VEWMOGLALLSEAVLGRQALLVNSQWPFLQHVDKAVSGSLSLTTLRAGAQEAIS 142

QY 121 PPDAAASAPLRTTADTPRKLFVYNSFLRGKLUYTGAEACTGD 165
 DB 143 PPDAAASAPLRTTADTPRKLFVYNSFLRGKLUYTGAEACTGD 187

RESULT 48

AAP8195
 ID AAP8195 standard; protein; 188 AA.

AC AAP8195;
 XX DT 25-MAR-2003 (revised)
 DT 20-NOV-1990 (first entry)

DB Erythropoietin encoded by EPO 140B.

XX EPO; erythropoietin; anaemia; renal failure.

XX OS Homo sapiens.

FH Key
 Peptide
 FT /label= leader sequence
 Protein
 FT /label= EPO

XX EP267678-A.

XX 18-MAY-1988.

XX 15-SEP-1987; 87EP-00308130.

XX 15-SEP-1986; 86US-00907369.

XX (INUA) INTEGRATED GENETICS INC.

PT Beck AK, Withy RM, Zabrecky JR, Masiello NC;

XX WPI; 1988-134531/20.

DR N-PSDB; AAN81554.

XX Recombinant human erythropoietin - produced by a transformed rodent
 PT epithelial cell capable of producing N-linked and O-linked glycosylated
 PT human erythropoietin.

PS Disclosure; Page ?; 23pp; English.

RESULT 49

ADFL6388
 ID ADF16388 standard; protein; 192 AA.

XX AC ADF16388;
 XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID1690.

XX KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human; gene; ds.

OS Homo sapiens.

XX PN WO2003060071-A2.

XX PD 24-JUN-2003.

XX PR 23-DEC-2002; 2002WO-US040891.

XX PR 21-DEC-2001; 2001US-0341811P.

PR 24-JAN-2002; 2002US-0350358P.

PR 28-JAN-2002; 2002US-0351360P.

PR 26-FEB-2002; 2002US-0359370P.

PR 28-FEB-2002; 2002US-036000P.

PR 27-MAR-2002; 2002US-036750P.

PR 08-APR-2002; 2002US-0370227P.

PR 10-MAY-2002; 2002US-037950P.

PR 24-MAY-2002; 2002US-038217P.

PR 28-MAY-2002; 2002US-0383123P.

PR 05-JUN-2002; 2002US-038570P.

PR 10-JUN-2002; 2002US-039425P.

PR 24-JUN-2002; 2002US-039800P.

PR 09-AUG-2002; 2002US-0402131P.

PR 13-AUG-2002; 2002US-040270P.

PR 18-SEP-2002; 2002US-041135P.

PR 18-SEP-2002; 2002US-0411426P.

PR 02-OCT-2002; 2002US-0414984P.

PR 11-OCT-2002; 2002US-0417611P.

PR 23-OCT-2002; 2002US-0420246P.

PR 05-NOV-2002; 2002US-0423623P.
 PR XX
 PA (HUMA-) HUMAN GENOME SCI INC.
 PA (DEBZ) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPIA PHARM CORP.
 XX
 PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX DR WPI; 2003-598517/56.
 DR N-PSDB; ADF16262.
 XX PT New albumin fusion protein, useful for preparing a composition for
 PT treating diabetes mellitus.
 XX PS Example 4; SEQ ID NO 1690; 24pp; English.
 XX CC This invention relates to a novel albumin fusion protein having albumin
 or biological activity. Human serum albumin is responsible for a
 significant proportion of the osmotic pressure of serum and also
 functions as a carrier of endogenous and exogenous ligands. The fusion of
 albumin to a therapeutic protein may increase shelf-life and stability of
 the therapeutic protein. The albumin fusion protein of the invention may
 allow production of compositions with antidiabetic activity whilst the
 nucleotide sequence which encodes it may be useful for gene therapy. The
 albumin fusion protein is useful for preparing a composition for treating
 diabetes mellitus. The present sequence is that of a therapeutic protein
 which was fused with human albumin to create a novel albumin fusion
 protein of the invention. Note: The sequence data for this patent did not
 form part of the printed specification, but was obtained in electronic
 format directly from WIPO at ftp://wipo.int/pub/publishedpat_sequences
 CC XX
 CC Sequence 192 AA;
 SQ
 Query Match 100.0%; Score 846; DB 7; Length 192;
 Best Local Similarity 100.0%; Pred. No. 2.7e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC OY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWNEVGQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWNEVGQA 87
 CC QY 61 VEVNQGLALLSEAVRGQALVNNSQWPWPLQLHQVDAVSGRLSLTTLRALGQKEAIS 120
 Db 88 VEVNQGLALLSEAVRGQALVNNSQWPWPLQLHQVDAVSGRLSLTTLRALGQKEAIS 147
 CC QY 121 PPDASAAAPRTTADTRKLFRVYSNPLRGKLYTGEACRTGD 165
 Db 148 PPDASAAAPRTTADTRKLFRVYSNPLRGKLYTGEACRTGD 192
 CC XX
 RESULT 50
 ADF16589
 ID ADF16589 standard; protein; 192 AA.
 XX AC ADF16589;
 XX DT 12-FEB-2004 (first entry)
 XX DB Human albumin fusion protein-related protein SeqID1691.
 XX KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human; gene; ds.
 XX OS Homo sapiens.
 XX PN WO2003050071-A2.
 XX PD 24-JUL-2003.
 XX PP 23-DEC-2002; 2002WO-US040891.
 XX PR 21-DEC-2001; 2001US-0341811P.
 XX
 PR 24-JAN-2002; 2002US-0350358P.
 PR 28-JAN-2002; 2002US-035130P.
 PR 26-FEB-2002; 2002US-0359370P.
 PR 28-FEB-2002; 2002US-036000P.
 PR 27-MAR-2002; 2002US-036750P.
 PR 08-APR-2002; 2002US-0370227P.
 PR 10-MAY-2002; 2002US-037890P.
 PR 24-MAY-2002; 2002US-0382617P.
 PR 28-MAY-2002; 2002US-0383123P.
 PR 05-JUN-2002; 2002US-038578P.
 PR 10-JUL-2002; 2002US-0394625P.
 PR 24-JUL-2002; 2002US-039808P.
 PR 09-AUG-2002; 2002US-0402131P.
 PR 13-AUG-2002; 2002US-0402708P.
 PR 18-SEP-2002; 2002US-041135P.
 PR 03-OCT-2002; 2002US-041494P.
 PR 23-OCT-2002; 2002US-042026P.
 PR 05-NOV-2002; 2002US-0423623P.
 XX PA (HUMA-) HUMAN GENOME SCI INC.
 PA (DEBZ) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPIA PHARM CORP.
 XX PR XX
 PR Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX DR WPI; 2003-598517/56.
 DR N-PSDB; ADF16263.
 XX PT New albumin fusion protein, useful for preparing a composition for
 PT treating diabetes mellitus.
 XX PS Example 4; SEQ ID NO 1691; 24pp; English.
 XX CC This invention relates to a novel albumin fusion protein having albumin
 or biological activity. Human serum albumin is responsible for a
 significant proportion of the osmotic pressure of serum and also
 functions as a carrier of endogenous and exogenous ligands. The fusion of
 albumin to a therapeutic protein may increase shelf-life and stability of
 the therapeutic protein. The albumin fusion protein of the invention may
 allow production of compositions with antidiabetic activity whilst the
 nucleotide sequence which encodes it may be useful for gene therapy. The
 albumin fusion protein is useful for preparing a composition for treating
 diabetes mellitus. The present sequence is that of a therapeutic protein
 which was fused with human albumin to create a novel albumin fusion
 protein of the invention. Note: The sequence data for this patent did not
 form part of the printed specification, but was obtained in electronic
 format directly from WIPO at ftp://wipo.int/pub/publishedpat_sequences
 CC XX
 CC Sequence 192 AA;
 SQ
 Query Match 100.0%; Score 846; DB 7; Length 192;
 Best Local Similarity 100.0%; Pred. No. 2.7e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC OY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWNEVGQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWNEVGQA 87
 CC QY 61 VEVNQGLALLSEAVRGQALVNNSQWPWPLQLHQVDAVSGRLSLTTLRALGQKEAIS 120
 Db 88 VEVNQGLALLSEAVRGQALVNNSQWPWPLQLHQVDAVSGRLSLTTLRALGQKEAIS 147
 CC QY 121 PPDASAAAPRTTADTRKLFRVYSNPLRGKLYTGEACRTGD 165
 Db 148 PPDASAAAPRTTADTRKLFRVYSNPLRGKLYTGEACRTGD 192
 CC XX
 RESULT 51
 ADF15305
 ID ADF15305 standard; protein; 192 AA.
 XX

This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is that of a therapeutic protein which was fused with human albumin to create a novel albumin fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/publishedpat_sequences

SQ Sequence 192 AA;

Query Match 100.0%; Score 846; DB 7; Length 192;
Best Local Similarity 100.0%; Pred. No. 2.7e-86; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative 0; MisMatches 0;

QY 1 APPRLICDSRVRLYLEAKEAENITTCGAECHSLENNTIVPTKVNFTYAWKMEVQQA 60
Db 28 APPRLICDSRVRLYLEAKEAENITTCGAECHSLENNTIVPTKVNFTYAWKMEVQQA 87
QY 61 VEWQGLALISEAVLRLQGALLVNSQPEPLQLHVDAVGSLSLTIRALGAQEAIIS 120
Db 88 VEWQGLALISEAVLRLQGALLVNSQPEPLQLHVDAVGSLSLTIRALGAQEAIIS 147
QY 121 PPDRASAAPLRTTADTRKLFRVYSNFLRGKLUYTGEACRGD 165
Db 148 PPDRASAAPLRTTADTRKLFRVYSNFLRGKLUYTGEACRGD 192

RESULT 53

ID ADF16726
ID ADF16726 standard; protein; 192 AA.

AC ADF16726;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID192.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

XX ID ADF16726
ID ADF16726 standard; protein; 192 AA.

AC ADF16726;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID192.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

XX ID ADF15296
ID ADF15296 standard; protein; 192 AA.

AC ADF15296;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID594.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

XX ID ADF15296
ID ADF15296 standard; protein; 192 AA.

AC ADF15296;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID594.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

PR 02-OCT-2002; 2002US-041494P.
PR 11-OCT-2002; 2002US-041761P.
PR 23-OCT-2002; 2002US-0420246P.
PR 05-NOV-2002; 2002US-0423623P.
XX PA (HUMA-) HUMAN GENOME SCI INC.
(DELB-) DELTA BIOTECHNOLOGY LTD.
(PRIN-) PRINCIPIA PHARM CORP.

XX PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
DR DR N-PSDB; ADF16400.

XX PT New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.

XX PS Example 4; SEQ ID NO 1828; 24PP; English.

This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is that of a therapeutic protein which was fused with human albumin to create a novel albumin fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/publishedpat_sequences

SQ Sequence 192 AA;

Query Match 100.0%; Score 846; DB 7; Length 192;
Best Local Similarity 100.0%; Pred. No. 2.7e-86; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative 0; MisMatches 0;

QY 1 APPRLICDSRVRLYLEAKEAENITTCGAECHSLENNTIVPTKVNFTYAWKMEVQQA 60
Db 28 APPRLICDSRVRLYLEAKEAENITTCGAECHSLENNTIVPTKVNFTYAWKMEVQQA 87
QY 61 VEWQGLALISEAVLRLQGALLVNSQPEPLQLHVDAVGSLSLTIRALGAQEAIIS 120
Db 88 VEWQGLALISEAVLRLQGALLVNSQPEPLQLHVDAVGSLSLTIRALGAQEAIIS 147
QY 121 PPDRASAAPLRTTADTRKLFRVYSNFLRGKLUYTGEACRGD 165
Db 148 PPDRASAAPLRTTADTRKLFRVYSNFLRGKLUYTGEACRGD 192

RESULT 54

ID ADF15296

AC ADF15296;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID594.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

XX ID ADF15296
ID ADF15296 standard; protein; 192 AA.

AC ADF15296;

XX DT 12-FEB-2004 (first entry)

DB Human albumin fusion protein-related protein SeqID594.

XX KW albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human; gene; db.

XX OS Homo sapiens.

PN WO2003060071-A2.

XX

PD 24-JUL-2003.

PR	23-DEC-2002;	2002WO-US040891.							
DR	21-DEC-2001;	2001US-0341811P.							
PR	24-JAN-2002;	2002US-0350358P.							
PR	28-JAN-2002;	2002US-0351360P.							
PR	26-FEB-2002;	2002US-0359370P.							
PR	28-FEB-2002;	2002US-036000P.							
PR	27-MAR-2002;	2002US-0367500P.							
PR	08-APR-2002;	2002US-0370227P.							
PR	10-MAY-2002;	2002US-0378950P.							
PR	24-MAY-2002;	2002US-0382617P.							
PR	28-MAY-2002;	2002US-0383123P.							
PR	05-JUN-2002;	2002US-0385708P.							
PR	10-JUL-2002;	2002US-0394625P.							
PR	24-JUL-2002;	2002US-0398008P.							
PR	09-AUG-2002;	2002US-0402131P.							
PR	13-AUG-2002;	2002US-0402708P.							
PR	18-SEP-2002;	2002US-0411426P.							
PR	02-OCT-2002;	2002US-0414984P.							
PR	11-OCT-2002;	2002US-0417611P.							
PR	23-OCT-2002;	2002US-0420246P.							
PR	05-NOV-2002;	2002US-0423623P.							
XX									
PA	(HUMA-)	HUMAN GENOME SCI INC.							
PA	(DELB)	DELTA BIOTECHNOLOGY LTD.							
PA	(PRIN-)	PRINCIPIA PHARM CORP.							
XX									
PT	Ballance DJ,	Turner AJ,	Rosen CA,	Haseltine WA;					
DR	WPI;	2003-598517/56.							
DR	N-PSDB;	ADF15861.							
XX									
PT	New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.								
XX									
PS	Example 4;	SEQ ID NO 594;	24pp;	English.					
XX									
CC	This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is that of a therapeutic protein which was fused with human albumin to create a novel albumin fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/published/pct_sequences								
XX									
SQ	Sequence 192 AA;								
XX									
Query Match	100.0%	Score 846;	DB 7;	Length 192;					
Best Local Similarity	100.0%	Score 846;	DB 7;	Length 192;					
Matches	165;	Conservative	0;	Mismatches	0;	Indels	0;	Gaps	0;
Qy	1 APPRLICDSRVLRVLYLEAKAENTITGCAEHCSLENENTVDPDTKVNFKWKRMEVGQAA	60							
Db	28 APPRLICDSRVLRVLYLEAKAENTITGCAEHCSLENENTVDPDTKVNFKWKRMEVGQAA	87							
Qy	61 VETWQGLALLSEAVTRQQLVNSSQMPQLQHDKAVSGLRITTLRAGQKAIS	120							
Db	88 VETWQGLALLSEAVTRQQLVNSSQMPQLQHDKAVSGLRITTLRAGQKAIS	147							
Qy	121 PPDDASAAPLRTTADTFRKLFYRSVNFRLGKLYKTYGACRTGD	165							
Db	148 PPDDASAAPLRTTADTFRKLFYTSNPLRGKLYKTYGACRTGD	192							
XX									
PT	New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.								
XX									
PS	Example 4;	SEQ ID NO 1830;	24pp;	English.					
XX									
CC	This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is that of a therapeutic protein which was fused with human albumin to create a novel albumin fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic								

Query Match Score 846; DB 7; Length 192;
 Best Local Similarity 100.0%; Pred. No. 2.7e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRLLICDSRVLYRVLLEAKAENITTCGAECNSLENITVTDKVNHYAWKMEVGQA 60
 DB 28 APPRLICDSRVLYRVLLEAKAENITTCGAECNSLENITVTDKVNHYAWKMEVGQA 87
 QY 61 VEWQGALLSEAVLVRGQALLVNSQWEPFLQHVDAVSGRSLSLTIRALGAQEAIIS 120
 DB 88 VEWQGALLSEAVLVRGQALLVNSQWEPFLQHVDAVSGRSLSLTIRALGAQEAIIS 147
 121 PPDASAAAPLRTTADTRKLFRVYSHNPLRGKLUYGEACTRGD 165
 148 PPDASAAAPLRTTADTRKLFRVYSHNPLRGKLUYGEACTRGD 192

RESULT 56
 ADF15295
 ID ADF15295 standard; protein; 192 AA.
 XX
 AC ADF15295;
 XX DT 12-FEB-2004 (first entry)
 XX Human albumin fusion protein-related protein SeqID593.
 KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human; gene; ds.
 OS Homo sapiens.
 XX PN WO2003060071-A2.
 PD 24-JUL-2003.
 XX
 PR 23-DEC-2002; 2002WO-US040891.
 XX
 PR 21-DEC-2001; 2001US-0341811P.
 PR 24-JAN-2002; 2002US-0350358P.
 PR 28-JAN-2002; 2002US-051360P.
 PR 26-FEB-2002; 2002US-0359370P.
 PR 28-FEB-2002; 2002US-0360000P.
 PR 27-MAR-2002; 2002US-0367500P.
 PR 08-APR-2002; 2002US-0370277P.
 PR 10-MAY-2002; 2002US-038950P.
 PR 24-MAY-2002; 2002US-0382617P.
 PR 28-MAY-2002; 2002US-0393123P.
 PR 05-JUN-2002; 2002US-0385708P.
 PR 10-JUL-2002; 2002US-0394625P.
 PR 24-JUL-2002; 2002US-0398008P.
 PR 09-AUG-2002; 2002US-0402131P.
 PR 13-AUG-2002; 2002US-0402708P.
 PR 18-SEP-2002; 2002US-0411355P.
 PR 02-OCT-2002; 2002US-0411426P.
 PR 11-OCT-2002; 2002US-0417611P.
 PR 23-OCT-2002; 2002US-0420246P.
 PR 05-NOV-2002; 2002US-0423623P.
 PR XX
 PA (HOMA-) HUMAN GENOME SCI INC.
 PA (DELTZ-) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPIA PHARM CORP.
 XX
 PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX
 DR WPI: 2003-598517/56.
 DR N-PSDB; ADF15860.
 XX
 PT New albumin fusion protein, useful for preparing a composition for
 treating diabetes mellitus.

RESULT 56
 ADF15295
 ID ADF15295 standard; protein; 192 AA.
 XX
 AC ADF15295;
 XX DT 12-FEB-2004 (first entry)
 XX Human albumin fusion protein-related protein SeqID593.
 KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human; gene; ds.
 OS Homo sapiens.
 XX PN WO2003060071-A2.
 PD 24-JUL-2003.
 XX
 PR 23-DEC-2002; 2002WO-US040891.
 XX
 PR 21-DEC-2001; 2001US-0341811P.
 PR 24-JAN-2002; 2002US-0350358P.
 PR 28-JAN-2002; 2002US-0351360P.
 PR 24-FEB-2002; 2002US-0359370P.
 PR 28-FEB-2002; 2002US-0360000P.
 PR 27-MAR-2002; 2002US-0367500P.
 PR 08-APR-2002; 2002US-0370277P.
 PR 12-FEB-2004 (first entry)
 XX Human albumin fusion protein-related protein SeqID1689.
 KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human; gene; ds.
 OS Homo sapiens.
 XX PN WO2003060071-A2.
 PD 24-JUL-2003.
 XX
 PR 23-DEC-2002; 2002WO-US040891.
 XX
 PR 21-DEC-2001; 2001US-0341811P.
 PR 24-JAN-2002; 2002US-0350358P.
 PR 28-JAN-2002; 2002US-0351360P.
 PR 24-FEB-2002; 2002US-0359370P.
 PR 28-FEB-2002; 2002US-0360000P.
 PR 27-MAR-2002; 2002US-0367500P.
 PR 08-APR-2002; 2002US-0370277P.
 PR 10-MAY-2002; 2002US-0378950P.
 PR 24-MAY-2002; 2002US-0388617P.
 PR 28-MAY-2002; 2002US-0383123P.
 PR 05-JUN-2002; 2002US-0385708P.
 PR 10-JUL-2002; 2002US-0394625P.
 PR 24-JUL-2002; 2002US-0398008P.
 PR 09-AUG-2002; 2002US-0402131P.

XX
 PS Example 4; SEQ ID NO 593; 24pp; English.
 XX
 CC This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the oncotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is that of a therapeutic protein which was fused with human albumin to create a novel albumin fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/publishedpat_sequences

PR 13-AUG-2002; 2002US-0402708P.
 PR 18-SEP-2002; 2002US-0411355P.
 PR 18-SEP-2002; 2002US-0411426P.
 PR 01-OCT-2002; 2002US-0414984P.
 PR 11-OCT-2002; 2002US-0417611P.
 PR 23-OCT-2002; 2002US-0420246P.
 PR 05-NOV-2002; 2002US-0423623P.
 XX (HOMA-) HUMAN GENOME SCI. INC.
 PA (DELLZ) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPA PHARM CORP.
 XX Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 PT DR WPI; 2003-599517/56.
 XX DR N-PSDB; ADF16261.
 PT New albumin fusion protein, useful for preparing a composition for
 PT treating diabetes mellitus.
 XX PS Example 4; SEQ ID NO 1689; 24pp; English.
 XX CC This invention relates to a novel albumin fusion protein having albumin
 CC or biological activity. Human serum albumin is responsible for a
 CC significant proportion of the osmotic pressure of serum and also
 CC functions as a carrier of endogenous and exogenous ligands. The fusion of
 CC albumin to a therapeutic protein may increase shelf-life and stability of
 CC the therapeutic protein. The albumin fusion protein of the invention may
 CC allow production of compositions with antidiabetic activity whilst the
 CC nucleotide sequence which encodes it may be useful for gene therapy. The
 CC albumin fusion protein is useful for preparing a composition for treating
 CC diabetes mellitus. The present sequence is that of a therapeutic protein
 CC which was fused with human albumin to create a novel albumin fusion
 CC protein of the invention. Note: The sequence data for this patent did not
 CC form part of the printed specification, but was obtained in electronic
 CC format directly from WIPO at ftp.wipo.int/pub/publishedpat_sequences
 XX SQ Sequence 192 AA;
 Query Match 100.0%; Score 846; DB 7; Length 192;
 Best Local Similarity 100.0%; Pred. No. 2.7e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC QY 1 APPRLICDSVLERVLLEAKEAENTTGCACHECSLENNTIPDTKVNFIYAWGRMEVGQAA 60
 Db 28 APPRLICDSVLERVLLEAKEAENTTGCACHECSLENNTIPDTKVNFIYAWGRMEVGQAA 87
 28 APPRLICDSVLERVLLEAKEAENTTGCACHECSLENNTIPDTKVNFIYAWGRMEVGQAA 87
 CC QY 61 VEWVQGLALLISEAVIRGQALLVNSQWPWELQLHDYDKAWSGLRSITTLRAGQKEAIS 120
 Db 88 VEWVQGLALLISEAVIRGQALLVNSQWPWELQLHDYDKAWSGLRSITTLRAGQKEAIS 147
 88 VEWVQGLALLISEAVIRGQALLVNSQWPWELQLHDYDKAWSGLRSITTLRAGQKEAIS 147
 CC QY 121 PPDAAASAAPLRTITADTRFLKFRVYNSFLRGKLYTGCAECACTGD 165
 Db 148 PPDAAASAAPLRTITADTRFLKFRVYNSFLRGKLYTGCAECACTGD 192
 148 PPDAAASAAPLRTITADTRFLKFRVYNSFLRGKLYTGCAECACTGD 192
 RESULT 58
 AAP50300 ID AAP50300 standard; protein; 193 AA.
 XX AC AAP50300;
 XX DT 25-MAR-2003 (revised)
 DT 01-JAN-1980 (first entry)
 DB Human erythropoietin encoded by positive clone (phage lambda-hE1) isolated
 DB from human fetal liver gene bank.
 KW Erythropoietin; red blood cell; erythrocyte; anaemia; blood; disorder;
 KW ss; phage lambda-hE1; gene bank.
 OS Homo sapiens.
 XX PN 03-DEC-1985; 85MO-US002405.
 XX PN WO8603520-A.
 XX PD 19-JUN-1986.
 XX PR 04-DEC-1984; 84US-00677813.
 PR 03-JAN-1985; 85US-0068622.
 PR 22-JAN-1985; 85US-0063258.
 PN WO8502610-A.
 XX PD 20-JUN-1985.
 XX PR 11-DEC-1984; 84WO-US002021.
 PR 13-DEC-1983; 83US-00501024.
 PR 21-FEB-1984; 84US-00502185.
 PR 28-SEP-1984; 84US-0055841.
 PR 30-NOV-1984; 84US-00675298.
 XX PA (KIRI) KIRIN AMGEN INC.
 XX DR WPI; 1983-159229/26.
 XX DR N-PSDB; AAN50347.
 PT New polypeptide having properties of erythropoietin - is prep'd. by
 PT cultivation of transformed eucaryotic or procaryotic host.
 XX PS Disclosure; Page 43; 113pp; English.
 XX CC Human erythropoietin encoded by a sequence encoded by this phage lambda-
 CC hel is essential for red blood cell formation and is used for the
 CC diagnosis and treatment of blood disorders such as anaemia. Large amounts
 CC of EPO may be obtained using recombinant DNA techniques in contrast to
 CC small amounts obtained from plasma and urine. This sequence is expressed
 CC in E. coli. See also AAN50345-6, AAN50348-50 and AAP50298-99, AAP50301.
 CC (Updated on 25-MAR-2003 to correct PA field.)
 XX SQ Sequence 193 AA;
 Query Match 100.0%; Score 846; DB 1; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC QY 1 APPRLICDSVLERVLLEAKEAENTTGCACHECSLENNTIPDTKVNFIYAWGRMEVGQAA 60
 Db 28 APPRLICDSVLERVLLEAKEAENTTGCACHECSLENNTIPDTKVNFIYAWGRMEVGQAA 87
 61 VEWVQGLALLISEAVIRGQALLVNSQWPWELQLHDYDKAWSGLRSITTLRAGQKEAIS 120
 88 VEWVQGLALLISEAVIRGQALLVNSQWPWELQLHDYDKAWSGLRSITTLRAGQKEAIS 147
 CC QY 121 PPDAAASAAPLRTITADTRFLKFRVYNSFLRGKLYTGCAECACTGD 165
 Db 148 PPDAAASAAPLRTITADTRFLKFRVYNSFLRGKLYTGCAECACTGD 192
 RESULT 59
 AAP50597 ID AAP50597 standard; protein; 193 AA.
 XX AC AAP50597;
 XX DT 25-MAR-2003 (revised)
 DT 01-JAN-1980 (first entry)
 XX DE Clone lambda HBPOFL13 encoding human erythropoietin.
 XX KW Erythropoietin; lambda HBPOFL13; recombinant plasmid vector; anaemia;
 KW mammal cell culture; 3T3; CHO; Chinese hamster ovary; BB.
 XX OS Homo sapiens.
 XX PN WO8603520-A.
 XX PD 19-JUN-1986.
 XX PR 03-DEC-1985; 85MO-US002405.
 XX PN WO8603520-A.
 XX PD 19-JUN-1986.
 XX PR 04-DEC-1984; 84US-00677813.
 PR 03-JAN-1985; 85US-0068622.
 PR 22-JAN-1985; 85US-0063258.
 XX OS Homo sapiens.

XX
PA
(FRIT/) GENETICS INST INC.
PA
Fritsch E, Hewick RM, Jacobs K,
XX
DR WPI; 1986-169459/26.
N-PSDB; AAN60513.

PT Prod. of human cDNA clone expressing erythropoietin - for mass prodn. of
XX
erythropoietin, useful for treating anaemia.
PS Disclosure; Page 7; 61pp; English.

CC A recombinant plasmid vector expressing this clone is expressed in e. g.
3T3 or CHO cell cultures. The produced erythropoietin is useful for
treatment of anaemia, especially renal anaemia. The cloned gene expresses
high levels of the protein and thus provides a means of mass production.
See also AAN60514-21 and AAN60598-99. (Updated on 25-MAR-2003 to correct
PA field.)

CC Sequence 193 AA;

XX SQ

Query Match	100.0%	Score 846;	DB 1;	Length 193;
Best Local Similarity	100.0%	Pred. No.	2. 8e-86;	
Matches	165;	Mismatches	0;	Indels 0;
				Gaps 0;

QY 1 APPRLICDSRVLYLEAKEAENITGCAEHCSLENNTVDPDKUNFYAWKMEVGQA 60

Db 28 APPRLICDSRVLYLEAKEAENITGCAEHCSLENNTVDPDKUNFYAWKMEVGQA 87

QY 61 VEWVOGLALLSEAVLRQQLVNSQWPMLQLHVDAVSGLRSLTTLRALGAQEAIIS 120

Db 88 VEWVOGLALLSEAVLRQQLVNSQWPMLQLHVDAVSGLRSLTTLRALGAQEAIIS 147

QY 121 PPDASAAAPLRTTADTPRLFLRIVSNFLRGKLUYTGEACRTGD 165

Db 148 PPDASAAAPLRTTADTPRLFLRIVSNFLRGKLUYTGEACRTGD 192

RESULT 60

AAP70256 ID AAB70256 standard; protein; 193 AA.

XX AC

XX DT 19-FEB-1991 (first entry)

XX OS Sequence of human erythropoietin (EPO).

XX KW Renal anaemia therapy; hormone.

XX OS Homo sapiens.

XX PH Key

FT Peptide 1. .27

FT /label= SIGNAL

FT Protein 28. .193

FT Region 81. .97

PT /notes "Fragment that Probe AAN70361 is based on"

XX PN EP232034-A.

XX PD 12-AUG-1987.

XX PP 19-JAN-1987; 87EP-00300399.

XX PR 23-JAN-1986; 86JP-0001286B.

XX PA (SUMO) SUMITOMO CHEM IND KK.

PA (SUMI-) SUMITOMI SEIYAKU KK.

XX PI Yanagi H, Ogawa I, Okamoto M, Hozumi T, Soga A, Yoshima T;

PI Teutsumi M;

XX DR WPI; 1987-223005/32.

DR N-PSDB; AAN70360, AAN70361.

XX PT Human erythropoietin prodn. - by culturing human cells; esp. Namalwa cells, transformed with DNA encoding human erythropoietin.

XX PS Disclosure; Fig 1; 22pp; English.

CC A cDNA library was prep. from the poly (A) RNA, which was isolated from the erythropoietin-producing human hepatoma cell Hp-1. The cDNA library was screened using the probes given in AAN70361 and AAN70362. A plasmid (named as p8-A20) was isolated. The nucleotide sequence of the cDNA obtained from this clone is shown in AAN70360

XX SQ Sequence 193 AA;

Query Match

100.0%	Score 846;	DB 1;	Length 193;	
Best Local Similarity	100.0%	Pred. No.	2. 8e-86;	
Matches	165;	Mismatches	0;	Indels 0;
				Gaps 0;

QY 1 APPRLICDSRVLYLEAKEAENITGCAEHCSLENNTVDPDKUNFYAWKMEVGQA 60

Db 28 APPRLICDSRVLYLEAKEAENITGCAEHCSLENNTVDPDKUNFYAWKMEVGQA 87

QY 61 VEWVOGLALLSEAVLRQQLVNSQWPMLQLHVDAVSGLRSLTTLRALGAQEAIIS 120

Db 88 VEWVOGLALLSEAVLRQQLVNSQWPMLQLHVDAVSGLRSLTTLRALGAQEAIIS 147

QY 121 PPDASAAAPLRTTADTPRLFLRIVSNFLRGKLUYTGEACRTGD 165

Db 148 PPDASAAAPLRTTADTPRLFLRIVSNFLRGKLUYTGEACRTGD 192

RESULT 61

AAR65499 ID AAR65499 standard; protein; 193 AA.

XX AC AAR65499;

XX DT 25-MAR-2003 (revised)

XX DE Human prepro-erythropoietin.

XX KW Erythropoietin; therapeutic; ss.

XX OS Synthetic.

XX PH Key

FT Peptide 1. .27

FT /notes "leader peptide"

XX PN WO9425055-A1.

XX PD 10-NOV-1994.

XX PP 29-APR-1994; 94WO-US004755.

XX PR 29-APR-1993; 93US-00055076.

XX PA (ABBO) ABBOTT LAB.

XX PI Okasinski GF, Devries PJ, Mellovitz BS, Meuth JL, Schaefer VG;

DR WPI; 1994-357906/44.

DR N-PSDB; AAQ74760.

XX PT Erythropoietin analogues - useful for treatment of anaemia and have enhanced erythropoietic effect.

XX PS Disclosure; Page 38-39; 55pp; English.

XX DNA encoding human prepro-erythropoietin may be ligated into an
 CC expression vector for erythropoietin expression in a CHO cell culture.
 CC Site-directed mutagenesis may be used in the construction of EPO
 CC analogues with improved activity, which may be used in pharmaceutical
 compositions for inducing erythropoiesis and treating anaemia. (Updated
 CC on 25-MAR-2003 to correct PN field.)
 XX SQ Sequence 193 AA;

Query Match	100.0%; Score 846; DB 2; Length 193;
Best Local Similarity	100.0%; Pred. No. 2.8e-86;
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAECISLNENITVPDTKVNFYAWKRMEVGQQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITTCGAECISLNENITVPDTKVNFYAWKRMEVGQQA 87

Qy 61 VEWQGGLLSEAVLRGQALLVNSQPEPLQLHVDKAVSGIRSLLTIRALGAQKRAIS 120
 Db 88 VEWQGGLLSEAVLRGQALLVNSQPEPLQLHVDKAVSGIRSLLTIRALGAQKRAIS 147

Qy 121 PPDASASAPLRTTADTPRKLFPRVYSNFLRGKLUKYGEACRTGD 165
 Db 148 PPDASASAPLRTTADTPRKLFPRVYSNFLRGKLUKYGEACRTGD 192

RESULT 62

AAR71137 ID AAR71137 standard; protein: 193 AA.

XX AC AAR71137;
 XX DT 17-OCT-1995 (revised)
 DE Human erythropoietin.

KW Human erythropoietin; glycosylation; salic acid; solubility; half-life;
 KW biological activity; proteolysis resistance; anaemia;
 KW chronic renal failure.

OS Homo sapiens.

Key Location/qualifiers
 FT Peptide 1..27
 FT /label= sig_peptide

WO9505465.A1.

PR 16-AUG-1994; 94WO-US009257.
 PR 17-AUG-1993; 93US-00108016.

PA (AMGE-) AMGEN INC.

XX PD 23-FEB-1995.

XX PI Elliott SG, Byrne TE;

XX DR WPI; 1995-194095/25.
 DR N-PSDB; AAQ92296.

XX PT Gene therapy for treatment of anaemia - and increasing red blood cell
 PT production by transforming red blood cells with the erythropoietin gene.
 XX Disclosure: Page 38-40; 51pp; English.

XX The amino acid sequence encoded by human EPO cDNA is given in AAR74141.
 CC Best Local Similarity 100.0%; Pred. No. 2.8e-86;
 CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC Sequence 193 AA;

CC the protein. The analogues are useful in claimed compns. for the
 CC treatment of chronic renal failure associated anaemia. (Updated on 25-MAR
 CC -2003 to correct PN field.)
 XX SQ Sequence 193 AA;

Query Match	100.0%; Score 846; DB 2; Length 193;
Best Local Similarity	100.0%; Pred. No. 2.8e-86;
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAECISLNENITVPDTKVNFYAWKRMEVGQQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITTCGAECISLNENITVPDTKVNFYAWKRMEVGQQA 87

Qy 61 VEWQGGLLSEAVLRGQALLVNSQPEPLQLHVDKAVSGIRSLLTIRALGAQKRAIS 120
 Db 88 VEWQGGLLSEAVLRGQALLVNSQPEPLQLHVDKAVSGIRSLLTIRALGAQKRAIS 147

Qy 121 PPDASASAPLRTTADTPRKLFPRVYSNFLRGKLUKYGEACRTGD 165
 Db 148 PPDASASAPLRTTADTPRKLFPRVYSNFLRGKLUKYGEACRTGD 192

RESULT 63

AAR74141 ID AAR74141 standard; protein: 193 AA.

XX AC AAR74141;
 XX DT 25-MAR-2003 (revised)
 DE 30-OCT-1995 (first entry)

XX Human erythropoietin.

KW Erythropoietin; anemia; gene therapy; gene transfer; red blood cell; RBC;
 KW erythrocyte; transformation; myoblast; EPO.

OS Homo sapiens.

XX PN WO9513376-A1.

XX DD 18-MAY-1995.

XX PP 09-NOV-1994; 94WO-US013066.

XX PR 10-NOV-1993; 93US-00149871.

PR 07-OCT-1994; 94US-00320480.

XX PA (AMGE-) AMGEN INC.
 PA (UYSC-) UNIV SOUTHERN CALIFORNIA.

XX PI Samal BB, Hamamori Y, Kedes LH;

XX DR WPI: 1995-194095/25.
 DR N-PSDB; AAQ92296.

XX PT Gene therapy for treatment of anaemia - and increasing red blood cell
 PT production by transforming red blood cells with the erythropoietin gene.
 XX Disclosure: Page 38-40; 51pp; English.

XX The amino acid sequence encoded by human EPO cDNA is given in AAR74141.
 CC Best Local Similarity 100.0%; Pred. No. 2.8e-86;
 CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC Sequence 193 AA;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTCGAECISLNENITVPDTKVNFYAWKRMEVGQQA 60

Db	28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	Db	28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA
Oy	61 VEWQGLALLSEAVLRLGQALLVSSQPEPLQLHDVKAVSGLSLTLLRAGKAIS	Oy	61 VEWQGLALLSEAVLRLGQALLVSSQPEPLQLHDVKAVSGLSLTLLRAGKAIS
Db	88 VEWQGLALLSEAVLRLGQALLVSSQPEPLQLHDVKAVSGLSLTLLRAGKAIS	Db	88 VEWQGLALLSEAVLRLGQALLVSSQPEPLQLHDVKAVSGLSLTLLRAGKAIS
Oy	121 PPDASAPLRTITADTPRKLFYVSNFLRGKLUYGEACRTGD 165	Oy	121 PPDASAPLRTITADTPRKLFYVSNFLRGKLUYGEACRTGD 165
Db	148 PPDASAPLRTITADTPRKLFYVSNFLRGKLUYGEACRTGD 192	Db	148 PPDASAPLRTITADTPRKLFYVSNFLRGKLUYGEACRTGD 192
RESULT 64			
AAR81982		AAR98397	
ID AAR81982	standard; protein; 193 AA.	ID AAR98397	standard; protein; 193 AA.
XX		XX	
AC AAR81982;		AC AAR98397;	
XX		XX	
DT 25-MAR-2003	(revised)	DT 15-SEP-1996	(first entry)
DT 27-FEB-1996		DE	
DB Human erythropoietin.		XX	
XX Erythropoietin; sialylation; sialic acid; glycosylation; reticulocyte; red blood cell; erythrocyte; haemocrit.		KW Erythropoietin; BPO; anaemia; gene therapy; vector; synthetic animal.	
XX Homo sapiens.		KW scaffold attachment region; SAR element; transgenic animal.	
OS		XX	
XX		OS Synthetic.	
FH Key	Location/Qualifiers	FH Key	Location/Qualifiers
FT Peptide	1.-27	FT Peptide	1.-27
FT /label= sig_peptide		FT /label= sig_peptide	
FT Modified-site	51 /label= N-glycosylation_site	FT Protein	28. .193
FT Modified-site	65 /label= N-glycosylation_site	FT Protein	/label= Mat_protein
FT Modified-site	110 /label= N-glycosylation_site	FT XX	
FT Modified-site	153 /label= O-glycosylation_site	PN W09619573-A1.	
FT		PN	
XX EP668351-A1.		PD 27-JUN-1996.	
XX PD 23-AUG-1995.		XX 18-DEC-1995; 95WO-GA000696.	
XX PF 12-OCT-1990;	95EP-00101849.	XX 19-DEC-1994; 94US-00358918.	
XX PR 13-OCT-1989;	89US-00421444.	XX PA (CANG-) CANGENE CORP.	
PR 09-OCT-1990;	90MC-US005758.	PA XX	
PA (AMCB-) AMGEN INC.		PI Delcuve G;	
XX Byrne TE, Elliott SG;		XX DR WPI; 1996-309587/31.	
XX DR N-PSDB; AAI31529, AAI31532.		XX DR N-PSDB; AAI31529, AAI31532.	
XX PS Claim 3; Page 58; 84pp; English.		XX Recombinant DNA molecule expressing mammalian erythropoietin - useful to transform cell lines, and for gene therapy, e.g. of anaemia's and other red blood cell disorders.	
XX PS		PT	
XX Disclosure; Fig 5; 31pp; English.		XX Claim 3; Page 58; 84pp; English.	
XX New human erythropoietin analogues with increased glycosylation - have increased activity useful for increasing prodn. of reticulocytes and red blood cells.		XX Human erythropoietin (BPO) (AAAR8397) functions to promote erythroid development, to initiate haemoglobin biosynthesis and to stimulate proliferation of immature erythroid precursors. It can be obt. by stable, long-term expression in mammalian cell lines transfected with a vector carrying BPO cDNA (AAAT31529) or genomic DNA (AAAT31532) operably linked to an expression control sequence and to 5' and 3' human apolipoprotein scaffold attachment region (SAR) elements (see also AAAT3150-31). Transgenic animals can be produced that express the recombinant BPO in their milk	
CC Human urinary erythropoietin (AAR81982) is a glycoprotein contg. 3 N-linked and 1 O-linked oligosaccharide chain. Erythropoietin analogues (AAR81982-87) have been produced in which the number of glycosylation sites is increased. (Updated on 25-MAR-2003 to correct PP field.)		CC Sequence 193 AA;	
CC SQ Sequence 193 AA;		CC Query Match 100.0%; Score 846; DB 2; Length 193;	
CC Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;		CC Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	
CC Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;		CC Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;	
Oy 1 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA		Oy 1 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	
Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA		Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	
Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA		Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	
Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA		Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	
Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA		Db 28 APPRLICDSRVLYRILLEKAEARNITTCGAECISLNENITVPTDKYFAYKRMEVQQA	

Query Match 100.0%; Score 846; DB 3; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2. 8e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 1 APPRICDSRIVERYLLEAKRAENITTGCAHCSLNENITYPDTKQFYAKRMEVGQQA 60
 DB 28 APPRLICDSRIVERYLLEAKRAENITTGCAHCSLNENITYPDTKQFYAKRMEVGQQA 87

QY 121 PPDASAAPLRTTADTRKLFRVYNSFLRGKLUKYGEACRTGD 165
 DB 88 VEWQGALLSEAVLHQALVNSSQPWEPLQHDKAVSGLSLTLLRAGQEAIS 120

QY 121 PPDASAAPLRTTADTRKLFRVYNSFLRGKLUKYGEACRTGD 165
 DB 88 VEWQGALLSEAVLHQALVNSSQPWEPLQHDKAVSGLSLTLLRAGQEAIS 120

RESULT 66
 ID AAY43398 standard; protein; 193 AA.
 XX AAY43398;
 AC
 XX
 DT 28-JAN-2000 (first entry)
 XX
 DB Human erythropoietin protein sequence.
 XX
 KW SAR element; scaffold attachment region; human; apolipoprotein B; tPA;
 tissue plasminogen activator; protein expression; Gene therapy; lysis;
 occlusive coronary artery thrombi; transmural myocardial infarction;
 KW ventricular function; congestive heart failure; acute ischaemic stroke;
 KW acute massive pulmonary embolism; venous thrombosis; arterial thrombosis;
 KW embolism; arteriovenous cannulae occlusion; plasminogen activator;
 KW intravenous catheter clearance; blood clot; erythropoietin.
 OS Homo sapiens.
 PN US5985607-A.
 XX
 PD 16-NOV-1999.
 XX
 PF 27-JUN-1997; 97US-00883795.
 XX
 PR 19-DEC-1994; 94US-00358918.
 XX
 PA (CANG-) CANGENE CORP.
 XX
 PT Awang G, Delcuve G;
 XX
 DR WPI; 2000-012788/01.
 DR N-FSDB; A2Z37201.
 XX
 PT Recombinant DNA molecules encoding tissue plasminogen activator proteins,
 operatively linked to a scaffold attachment region, useful for the
 production of tissue plasminogen activator both in vivo and in vitro.
 XX
 PS Example 2; Fig 3; 49pp; English.

This sequence represents the human erythropoietin protein. The invention relates to a recombinant DNA molecule adapted for expression of tissue plasminogen activator (tPA). The DNA molecule comprise a sequence encoding tPA, an expression control sequence operatively linked to the tPA sequence, and at least one human apolipoprotein B scaffold attachment region (SAR) element (the SAR is not a 5' proximal apolipoprotein B SAR). The SAR element is used to increase the expression of the coding sequences. The recombinant nucleic acids may be used for the recombinant production of tPA both in vitro or in vivo (e.g. as part of a gene therapy procedure). tPA may be administered to treat and remove blood clots. It is especially useful for the lysis of occlusive coronary artery thrombi associated with evolving transmural myocardial infarction to improve ventricular function and reduce the risk of congestive heart failure. Additionally, it may be used in the management of acute massive pulmonary embolism, venous thrombosis and acute ischaemic stroke. Finally, tPA may be used in treating arterial thrombosis or embolism, arteriovenous cannulae occlusion and intravenous catheter clearance. In contrast to other plasminogen activators (e.g. urokinase and streptokinase), the activity of tPA is relatively localised and (in theory) is less likely to produce systemic haemorrhagic disorders

XX Sequence 193 AA;

XX
 SQ

Query Match 100.0%; Score 846; DB 3; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2. 8e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRICDSRIVERYLLEAKRAENITTGCAHCSLNENITYPDTKQFYAKRMEVGQQA 60
 DB 28 APPRLICDSRIVERYLLEAKRAENITTGCAHCSLNENITYPDTKQFYAKRMEVGQQA 87

QY 121 PPDASAAPLRTTADTRKLFRVYNSFLRGKLUKYGEACRTGD 165
 DB 88 VEWQGALLSEAVLHQALVNSSQPWEPLQHDKAVSGLSLTLLRAGQEAIS 120

QY 121 PPDASAAPLRTTADTRKLFRVYNSFLRGKLUKYGEACRTGD 165
 DB 88 VEWQGALLSEAVLHQALVNSSQPWEPLQHDKAVSGLSLTLLRAGQEAIS 120

RESULT 67
 ID AAY94330
 ID AAY94330 standard; protein; 193 AA.
 XX
 AC AAY94330;
 XX
 DT 28-NOV-2000 (first entry)
 XX
 DB Human erythropoietin protein.
 XX
 KW Human; erythropoietin; Epo; glycosylation; anaemia; cancer; viral infection;
 KW chronic renal failure; myelosuppressive therapy; HIV; blood loss.
 XX
 OS Homo sapiens.
 XX
 PR 04-MAY-2000.
 XX
 RF 18-OCT-1999; 99WO-US024435.
 XX
 PR 23-OCT-1998; 98US-00178292.
 XX
 PA (AMGE-) AMGEN INC.
 XX
 PI Egrie JC, Elliott SG, Brown JK;
 XX
 DR WPI; 2000-350735/30.
 XX
 PT Raising and maintaining hematocrit in a mammal by administering an effective amount of a hyperglycosylated analog of erythropoietin, useful for treating anemia associated with myelosuppressive therapy or excessive blood loss.
 XX
 PS Disclosure; Fig 1; 63pp; English.

The present sequence is human erythropoietin (Epo). Epo is a glycoprotein hormone necessary for the maturation of erythroid progenitor cells into erythrocytes. It has been discovered that hyperglycosylated Epo has a longer half-life and greater in vivo activity than recombinant human Epo. Several hyperglycosylated Epo mutants (AAY94331 to AAY4544) have been made by in vitro mutagenesis. Hyperglycosylated Epo analogs are useful as they may be used instead of recombinant Epo to increase and maintain the level of red blood cells in mammals. The Epo analogs may be used to treat or prevent anaemia associated with chronic renal failure, myelosuppressive therapy, certain cancers, viral disease, such as HIV and excessive blood loss.

SQ	Sequence 193 AA;	Db	28 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA
Query Match	100.0%; Score 846; DB 3; Length 193;	QY	61 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 120
Best Local Similarity	100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	Db	88 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 147
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	QY	1 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 60
CC	28 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 87	Db	2 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 87
CC	61 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 120	QY	121 PPDASAAPLRTTADTRKURVYSFLRGKLYTGEACRTGD 165
CC	148 PPDASAAPLRTTADTRKURVYSFLRGKLYTGEACRTGD 192	Db	148 PPDASAAPLRTTADTRKURVYSFLRGKLYTGEACRTGD 192
SQ	Sequence 193 AA;	Db	RESULT 69
Query Match	100.0%; Score 846; DB 3; Length 193;	ID	AAV99704 standard; protein; 193 AA.
Best Local Similarity	100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	XX	AAV99704;
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	AC	AAV99704;
XX	25-SEP-2000 (first entry)	XX	15-SEP-2000 (first entry)
DE	Amino acid sequence of a human erythropoietin polypeptide.	DE	Human non-glycosylated erythropoietin NGE.
XX	Human; erythropoietin; EPO; inhibitor; nuclear factor-kappaB; NF-kappaB;	XX	Human; non-glycosylated erythropoietin; NGE; haemocrit; antianemic;
KW	multi-drug resistance gene; malignant hemopathy; solid tumour;	KW	anaemia; erythropoiesis promoter.
KW	malignant blood disease; leukaemia; lymphoma; solid cancer.	XX	Homo sapiens.
OS	Homo sapiens.	OS	Homo sapiens.
XX	WO200030587-A2.	XX	PN0002772-A2.
DD	02-JUN-2000.	DD	08-JUN-2000.
XX	24-NOV-1999; 99WO-FR002897.	XX	23-NOV-1999; 99WO-US027801.
PF	PR 25-NOV-1998; 98FR-00014858.	PF	30-NOV-1998; 98US-0110299P.
XX	(CNRS) CENT NAT RCH SCI.	XX	PA (ELLI) LILLY & CO ELI.
PA	XX	PI Beals JM, Glaesner W, Micanovic R, Milican RL, Witcher DR;	
XX	XX	DR WPI; 2000-412320/35.	
PR	XX	PT Non-glycosylated erythropoietic compound useful for increasing hematocrit level in mammal with insufficient hematocrit levels in conditions such as anaemia, comprises protein covalently bonded to polymer.	
XX	XX	PT Non-glycosylated erythropoietic compound useful for increasing hematocrit level in mammal with insufficient hematocrit levels in conditions such as anaemia, comprises protein covalently bonded to polymer.	
PR	XX	PS Claim 1; Page 91-92; 94pp; English.	
PI	XX	CC The present sequence is the non-glycosylated erythropoietin NGE. The protein promotes erythropoiesis and can therefore be used to increase haematocrit levels in mammal with conditions such as anaemia, in which levels of haematocrit are insufficient. Mutants derived from the present protein can also be used to treat such conditions. The analogues, designated NGEAs, do not themselves cause a significant increase in haematocrit but they acquire that property once they are derivatized with polyethylene glycol polymers. The analogues can be produced using a linkerless aldehyde modification process. They show stability and bioactivity in vivo. The compounds can be produced by recombinant DNA technology or by chemical procedures such as solution or solid-phase peptide synthesis	
PI	XX	CC Sequence 193 AA;	
XX	The present sequence represents a human erythropoietin (EPO) polypeptide. The human growth hormone protein is used as an inhibitor of the activation of nuclear factor kappaB (NF-kappaB). The inhibitor inhibits activation of NF-kappaB, and thus transcription of the multi-drug resistance gene (which contains binding sites for NF-kappaB within its regulatory regions). The inhibitors are used to produce pharmaceuticals which may be used in the treatment of malignant hemopathy or solid tumours. The inhibitors are especially used to treat malignant blood diseases (leukaemia, lymphoma) and solid cancers (of breast or ovary)	CC Query Match 100.0%; Score 846; DB 3; Length 193;	
PS	XX	CC Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	
N-PSDB	XX	CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	
DR	XX	CC SQ Sequence 193 AA;	
XX	The present sequence represents a human erythropoietin (EPO) polypeptide. The human growth hormone protein is used as an inhibitor of the activation of nuclear factor kappaB (NF-kappaB). The inhibitor inhibits activation of NF-kappaB, and thus transcription of the multi-drug resistance gene (which contains binding sites for NF-kappaB within its regulatory regions). The inhibitors are used to produce pharmaceuticals which may be used in the treatment of malignant hemopathy or solid tumours. The inhibitors are especially used to treat malignant blood diseases (leukaemia, lymphoma) and solid cancers (of breast or ovary)	CC Query Match 100.0%; Score 846; DB 3; Length 193;	
CC	XX	CC Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	
CC	CC Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	QY 1 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 60	
CC	CC	Db 2 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 87	
CC	CC	QY 61 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 120	
CC	CC	Db 88 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 147	
SQ	Sequence 193 AA;	Db	88 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 147
Query Match	100.0%; Score 846; DB 3; Length 193;	QY	1 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 60
Best Local Similarity	100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;	Db	2 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 87
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	QY	61 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 120
OY	1 APPRLICDSRVLERYLLEAKEAENITTCGAECRHSLENNTVDPDKVNYAWKMEVGQOA 60	Db	88 VEWQGLALLSEAVLRGQALLNSQWPWPLQLWDKAVSGRSLSLTILRAQKAIS 147

Qy 121 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165
 XX ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
 Db 148 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 192
 XX ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||

RESULT 70

ID AAB34978 standard; protein; 193 AA.

XX Human erythropoietin SEQ ID NO: 4.

XX DT 27-MAR-2001 (first entry)

XX DB Human erythropoietin; EPO; hybridisation probe; gene therapy;

XX KW Chimpanzee; erythropoietin; EPO; mapping; therapeutic agent.

XX OS Homo sapiens.

XX AC AAB34978; WO200068376-A1.

XX PD 16-NOV-2000.

XX PP 05-MAY-2000; 2000WO-US012370.

XX PR 07-MAY-1999; 99US-00307307.

XX PR 28-MAR-2000; 2000US-0287524P.

XX PR 19-APR-2000; 2000US-00552265.

XX PA (GETH) GENENTECH INC.

XX PI Desauvage F, Henner DJ,

XX DR 2001-007393/01.

XX PS Nucleic acids encoding chimpanzee erythropoietin, useful for treatment of e.g. anemia, also derived proteins, antibodies and modulators.

XX PS Disclosure; Fig 3; 09PP; English.

XX CC The present invention provides the coding and protein sequences of chimpanzee erythropoietin (EPO). These sequences can be used in gene therapy, to block the activity of EPO, as hybridisation probes, in genetic and chromosome mapping and as therapeutic agents

XX SQ sequence 193 AA:

Query Match 100.0%; Score 846; DB 4; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLYRLVLEAKEARNTTGCABCISLNENITVPDTKUNFYAWKRMVGQQA 60
 Db 28 APPRLICDSRVLYRLVLEAKEARNTTGCABCISLNENITVPDTKUNFYAWKRMVGQQA 87
 Qy 61 VEVWQGLALLSEAVLRGQALLNSQWEPLQHVDKAVSGLSLTLRAGQKAIS 120
 Db 88 VEVWQGLALLSEAVLRGQALLNSQWEPLQHVDKAVSGLSLTLRAGQKAIS 147
 Qy 121 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165
 Db 148 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 192
 RESULT 71
 ID AAB85573 standard; protein; 193 AA.

XX AC AAB85573;

XX DT 29-OCT-2001 (first entry)

XX Human erythropoietin (EPO) sequence.

XX DR /note= "mature protein"

XX KW Transgenic; pig; human; erythropoietin; EPO; milk; PMSG; hCG; chorionic gonadotrophic hormone; WAP promoter.

XX OS Homo sapiens.

XX Key /note= "signal peptide"

XX Peptide 1..27

XX FT Protein 28..193

XX FT /note= "mature protein"

XX DN WO200159074-A1.

XX PD 16-AUG-2001.

XX PP 28-JUN-2000; 2000WO-KR000675.

XX PR 14-FEB-2000; 2000KR-00006888.

XX PA (KORE-) REPUBLIC KOREA.

XX PI Chang W, Park J, Seong H, Min K, Yang B, Im G, Lee Y, Lee C;

XX PI Kim J,

XX DR N-PSDB; AAH46972.

XX PR Producing transgenic porcine that secretes human erythropoietin (hEPO) in milk, by introducing vector comprising hEPO genome into fertilized eggs of porcine to which PMSG and hCG were administered, and developing progeny.

XX PS Claim 4; Fig 3; 21PP; English.

CC The invention relates to producing transgenic pigs (P) that secrete human erythropoietin (hEPO) in milk. The method involves administering PMSG and human chorionic gonadotrophic hormone (hCG) into (P), collecting fertilized eggs after making, injecting expression vector containing a 2.6 kb WAP promoter, hEPO genome and SV40 poly A DNA into male pronuclei, transplanting them in surrogate mother pig and allowing it to give birth. The method provides transgenic porcine capable of secreting hEPO in their milk, thus producing the expensive useful medicine at a low cost with stability on a large scale, giving a contribution to the improvement of human health. The present sequence represents a human EPO sequence incorporated into the genome of porcine

XX SQ Sequence 193 AA:

Query Match 100.0%; Score 846; DB 4; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLYRLVLEAKEARNTTGCABCISLNENITVPDTKUNFYAWKRMVGQQA 60
 Db 28 APPRLICDSRVLYRLVLEAKEARNTTGCABCISLNENITVPDTKUNFYAWKRMVGQQA 87
 Qy 61 VEVWQGLALLSEAVLRGQALLNSQWEPLQHVDKAVSGLSLTLRAGQKAIS 120
 Db 88 VEVWQGLALLSEAVLRGQALLNSQWEPLQHVDKAVSGLSLTLRAGQKAIS 147
 Qy 121 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 165
 Db 148 PPDAAASAPLRTTADTPRKLFRVYNSFLRGKLUKYGEACRTGD 192
 RESULT 72
 ID AAB15341
 ID AAB15341 standard; protein; 193 AA.

XX AC AAB15341;

XX DT 29-OCT-2001 (first entry)

XX

PR 01-JUL-2002; 2002US-0392455P.

PR 03-JUL-2002; 2002US-0393423P.

XX

PA (WARR-) WARREN INST INC KENNETH S.

XX

PA (LUND) LUNDBECK AS H.

XX

PI Nielsen J, Pedersen JT, Gerwien J, Bay K, Pedersen LO, Leist M;

PI Geist MA, Kallunki P, Christensen S, Sager T, Brines M, Cerami A;

PI Cerami C;

XX

DR WPI; 2004-071985/07.

XX

PT New murein recombinant tissue protective cytokines and encoding nucleic acid molecules, useful for protecting, restoring or enhancing the viability of responsive cells, tissues or organs in mammals, including humans.

XX

PS Claim 4; SEQ ID NO 22; 323pp; English.

XX

The invention relates to a novel murein recombinant tissue protective cytokine lacking at least one activity selected from increasing haemocrit, vasoactive action, hyperactivating platelets, pro-coagulant activities and increasing production of thrombocytes. A murein of the invention has vasoactive, neuroprotective, nootropic, ophthalmological, cardiovascular, respiratory, nephrotopic, uropathic, gynaecological, gastrointestinal, and endocrine activity. A polynucleotide encoding a cytokine of the invention may have a use in gene therapy. The recombinant tissue protective cytokine is useful for preparing a pharmaceutical composition for the protection against and prevention of a tissue injury as well as the restoration of and rejuvenation of tissue and tissue function in a mammal, where the injury is caused by a seizure disorder, multiple sclerosis, stroke, hypotension, cardiac arrhythmia, myocardial infarction, inflammation, age-related loss of cognitive function, radiation damage, cerebral palsy, neurodegenerative disease, Alzheimer's disease, Parkinson's disease, Leigh disease, AIDS dementia, memory loss, amyotrophic lateral sclerosis, alcoholism, mood disorder, anxiety disorder, attention deficit disorder, autism, Creutzfeld-Jakob disease, brain or spinal cord trauma or ischaemia, heart-lung bypass, chronic heart failure, macular degeneration, diabetic neuropathy, diabetic retinopathy, glaucoma, retinal ischaemia, or retinal trauma. The composition and methods may be used for preventing or treating neurological disorders, opthalmic diseases, cardiovascular diseases, reproductive diseases, gastrointestinal diseases, kidney, urinary and metabolic abnormalities. The present sequence is used in the exemplification of the invention.

SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 8; Length 193; Best Local Similarity 100.0%; Pred. No. 2.8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLRVLYLEAKERENITGCAHCNSLENNTIPDTKVNFIYAKRMEVGQA 60
Db 28 APPRLICDSRVLRVLYLEAKERENITGCAHCNSLENNTIPDTKVNFIYAKRMEVGQA 87
QY 61 VEWMOGLALLSRAVLRGQALVNNSQWPPLQHDKAVSGRSLSLTLLRAIGAOKEAIIS 120
Db 88 VEWMOGLALLSRAVLRGQALVNNSQWPPLQHDKAVSGRSLSLTLLRAIGAOKEAIIS 147
Qy 121 PPDAAASAPLRTITADPFRKLVYSNPLRGKLKYGEACRTGD 165
Db 148 PPDAAASAPLRTITADPFRKLVYSNPLRGKLKYGEACRTGD 192

RESULT 78 ADL0601 ID ADL0601 standard; protein; 193 AA.
XX AC ADL0601;
XX DT 03-JUN-2004 (first entry)
XX Human 165 residue erythropoietin analogue #20.
XX Human; erythropoietin; EPO; iron distribution disturbance; diabetes;

DT 15-APR-2004 (First entry)

XX DE Human erythropoietin protein, SEQ ID No 108.

XX T-cell epitope; cytokine; receptor; CD4+; CD8+; immunogenicity; interferon-beta; tumour necrosis factor receptor-1; erythropoietin; thrombopoietin; inflammation; cancer; anaemia; human erythropoietin.

XX KW Homo sapiens.

XX PN WO2003104263-A2.

XX PD 18-DEC-2003.

XX PI Harding FA, Power SD;

XX DR 26-FEB-2003; 2003WO-US005917.

XX PR 01-MAY-2002; 2002US-0376743P.

XX PA (GEMV) GENENCOR INT INC.

XX PI

XX DR Determining T-cell epitope of a protein (e.g. cytokine or cytokine receptor), useful for reducing protein allergenicity, comprises combining PT differentiated dendritic cells and naive T-cells with a peptide having PT the T-cell epitope.

XX PS Claim 4; SEQ ID NO 108; 51pp; English.

XX The invention relates to a novel method for determining a T-cell epitope of a protein, where the protein is selected from cytokines and cytokine receptors. The method comprises combining a solution of differentiated dendritic cells and naive CD4+ and/or CD8+ T-cells with a peptide comprising the T-cell epitope. The composition and methods are useful in reducing the immunogenicity of cytokines and cytokine receptors such as interferon-beta, soluble tumour necrosis factor receptor-1, erythropoietin or thrombopoietin. These modified cytokines and cytokine receptors may be used for treating various conditions such as inflammation, cancer or anaemia. This sequence represents the human erythropoietin protein of the invention.

XX SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 8; Length 193; Best Local Similarity 100.0%; Pred. No. 2.8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLRVLYLEAKERENITGCAHCNSLENNTIPDTKVNFIYAKRMEVGQA 60
Db 28 APPRLICDSRVLRVLYLEAKERENITGCAHCNSLENNTIPDTKVNFIYAKRMEVGQA 87
QY 61 VEWMOGLALLSRAVLRGQALVNNSQWPPLQHDKAVSGRSLSLTLLRAIGAOKEAIIS 120
Db 88 VEWMOGLALLSRAVLRGQALVNNSQWPPLQHDKAVSGRSLSLTLLRAIGAOKEAIIS 147
Qy 121 PPDAAASAPLRTITADPFRKLVYSNPLRGKLKYGEACRTGD 165
Db 148 PPDAAASAPLRTITADPFRKLVYSNPLRGKLKYGEACRTGD 192

RESULT 79 ADL0601 ID ADL0601 standard; protein; 193 AA.
XX AC ADL0601;
XX DT 03-JUN-2004 (first entry)
XX Human 165 residue erythropoietin analogue #20.
XX Human; erythropoietin; EPO; iron distribution disturbance; diabetes;

KW non-insulin dependent diabetes; type 2 diabetes; reticulocyte production;
 KW red blood cell production; glycosylation site; analogue; antidiabetic;
 XX mutant; mutein.

OS Homo sapiens.

XX Synthetic.

XX WO2004019972-A1.

XX 11-MAR-2004.

XX 20-AUG-2003; 2003WO-EPO09194.

XX 29-AUG-2002; 2002EP-00019100.

XX PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX PI Lehmann P., Roeddiger R., Walter-Matsui R.

XX DR WPI; 2004-282643/26.

XX PT Use of erythropoietin protein in manufacture of medicament for treating disturbances of iron distribution in diabetes.

XX Disclosure; Page; 31pp; English.

XX The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in diabetes. The erythropoietin protein is preferably a human erythropoietin (e.g., epoetin alpha and epoetin beta) which may be expressed by endogenous gene activation or an erythropoietin analogue such as darbepoetin alpha. The erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation. Patients with diabetes have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effects of iron accumulation in certain organs, leading to organ damage and destruction, and/or experience effects similar to anaemia due to iron usage in blood cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticulocytes and red blood cells, and this has been found to have a beneficial effect on iron distribution disturbances in diabetes e.g., non-insulin dependent (type 2) diabetes. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron distribution in diabetes. Sequences ADL06782-ADL06806 represent analogues of the 165 amino acid human erythropoietin which contain additional or altered glycosylation sites.

Note: The present sequence is not shown in the specification, but is derived from the wild-type 165 residue human EPO (ADL06780) and the information given on page 6.

CC Note: The present sequence is not shown in the specification, but is derived from the wild-type 165 residue human EPO (ADL06780) and the information given on page 6.

XX Sequence 193 AA;

XX Query Match 100.0%; Score 846; DB 8; Length 193;

Best Local Similarity 100.0%; Pred. No. 2.8e-86; Length 193;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

CC QY 1 APPRICDSVRLERYLLEAKEAENITTGCAEHCSLENNTVPDTKVNFKWAKMEVGQAA 60

CC Db 1 APPRICDSVRLERYLLEAKEAENITTGCAEHCSLENNTVPDTKVNFKWAKMEVGQAA 60

CC QY 61 VEWVOGLALISEAVLRGQALLVNSQPWPQLQHDKAVSGLRSLTLLRGAQEAKIS 120

CC Db 121 PPDAAASAPRITATDPRFLFRVSNFLFGKLUYTGEACRTGD 165

CC 121 PPDAAASAPRITATDPRFLFRVSNFLFGKLUYTGEACRTGD 165

XX RESULT 80

ID AD059436 standard; protein; 193 AA.

XX AC AD059436;

XX DT 26-AUG-2004 (first entry)

XX DE Human 165 residue erythropoietin analogue #20.

XX Human; erythropoietin, EPO; iron distribution disturbance; heart disease; heart insufficiency; coronary heart disease; atherosclerosis; acute coronary syndrome; heart failure; congestive heart failure; reticulocyte production; red blood cell production; cardiac; antiatherosclerotic; glycosylation site; analogue; mutant; mutein.

XX OS Homo sapiens.

XX Synthetic.

XX PN WO2004047858-A1.

XX PR 10-JUN-2004.

XX PR 17-NOV-2003; 2003WO-EPO12822.

XX PR 22-NOV-2002; 2002EP-00026342.

XX PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX PI Lehmann P., Roeddiger R., Walter-Matsui R.

XX DR WPI; 2004-450212/42.

XX PT Use of erythropoietin protein in the manufacture of medicament for treating disturbances of iron distribution in heart diseases e.g. heart insufficiency.

XX Disclosure; Page; 31pp; English.

XX The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in heart diseases. The erythropoietin protein is preferably a human erythropoietin (e.g., epoetin alpha and epoetin beta) which may be expressed by endogenous gene activation or an erythropoietin analogue such as darbepoetin alpha. The erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation. Patients with heart diseases have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effects of iron accumulation in certain organs, leading to organ damage and destruction, and/or experience effects similar to anaemia due to iron usage in blood cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticulocytes and red blood cells, and this has been found to have a beneficial effect on iron distribution disturbances in heart diseases e.g., heart insufficiency, coronary heart disease, atherosclerosis, acute coronary syndrome, heart failure and congestive heart failure. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron distribution in heart diseases. Sequences AD059417-AD05941 represent analogues of the 165 amino acid human erythropoietin which contain additional or altered glycosylation sites. Note: The present sequence is not shown in the specification, but is derived from the wild-type 165 residue human EPO (AD059415) and the information given on page 6.

XX Sequence 193 AA;

XX Query Match 100.0%; Score 846; DB 8; Length 193;

Best Local Similarity 100.0%; Pred. No. 2.8e-86; Length 193;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

CC QY 1 APPRICDSVRLERYLLEAKEAENITTGCAEHCSLENNTVPDTKVNFKWAKMEVGQAA 60

CC Db 1 APPRICDSVRLERYLLEAKEAENITTGCAEHCSLENNTVPDTKVNFKWAKMEVGQAA 60

CC QY 61 VEWVOGLALISEAVLRGQALLVNSQPWPQLQHDKAVSGLRSLTLLRGAQEAKIS 120

Db 61 VEWQGALLSEAVURGQALLVNSQPWEPLQLHVDAVSLRSLTLLAIGAKEYAIS 120
 QY 121 PPDAAASAPLRTITADFRKLFPRVYSNPLRGKLKUYGAEACTGD 165
 Db 121 PPDAAASAPLRTITADFRKLFPRVYSNPLRGKLKUYGAEACTGD 165

RESULT 81

ID ADT07724 standard; protein; 193 AA.
 XX

AC ADT07724;

DT 13-JAN-2005 (first entry)

XX Human erythropoietin protein.

XX

KW Erythropoietin; EPO; reduced immunogenicity; reduced immunity;

KW major histocompatibility complex class II; MHC;

KW helper T lymphocyte response; HTL; fungal disease; viral disease;

KW bacterial disease; parasitic disease; cancer; autoimmune disease;

KW alloraft rejection; allergy; Lyme disease; ulcerative colitis;

KW transplantation; haemophilia; osteoporosis; metabolic disease;

KW food hypersensitivity; cytostatic; immunosuppressive; antiinflammatory;

KW human.

OS Homo sapiens.

XX WO2004089973-A2.

XX 21-OCT-2004.

PP 02-APR-2004; 2004WO-US010353.

XX PR 02-APR-2003; 2003US-0459939P.

(EPIM-) BIIMMUNE INC.

XX Tangri S, Mothe B, Sette A, Southwood S, Briggs K, Chestnut RW;

DR WPI; 2004-748719/73.

PT New isolated or purified modified erythropoietin construct useful for treatment of anemia comprises a sequence selected from 5 sequences each containing 193 amino acids as given in specification, or truncated modified erythropoietin.

PT XX

PS Example 1; SEQ ID NO 3; 223PP; English.

XX The invention relates to isolated or purified modified erythropoietin (EPO) constructs (MSC), and truncated modified erythropoietin constructs. These constructs are peptides, polypeptides, proteins or antibodies having reduced immunogenicity as compared to the naturally occurring form. Also disclosed is a method of producing such peptides. The reduced immunity is as a result of reduced binding to major histocompatibility complex (MHC) class II molecules. The peptides of the invention are useful for antagonizing the erythropoietin (EPO) receptor or treating diseases or conditions associated with over-activation of the EPO receptor. The invention is useful for producing a peptide, polypeptide, protein and antibody having reduced immunogenicity, which is useful in the treatment and diagnosis of diseases, conditions and disorders. It is also useful for reducing the helper T lymphocyte (HTL) response against a candidate protein. The peptides, polypeptides, proteins and antibodies are useful for the treatment of pathological states (such as fungal, viral, bacterial and parasitic diseases, cancer (such as breast cancer, non-Hodgkin's lymphoma), autoimmune diseases (such as rheumatoid arthritis, multiple sclerosis, myasthenia gravis), allograft rejection, allergies (e.g. pollen allergies), Lyme disease, hepatitis B and C, ICMV, post-streptococcal endocarditis or glomerulonephritis, ulcerative colitis, Crohn's disease, psoriasis, chronic renal failure, asthma, transplantation, haemophilia, Paget's disease, osteoporosis, chronic granulomatous disease, genital warts, diabetes, defective tissue growth,

CC metabolic disease and food hypersensitivities). The peptides, polypeptides, proteins and antibodies are modified so as to have reduced immunogenicity as a result of reduced binding to MHC class II against various DR and DQ molecules and the subsequent reduced helper T lymphocyte (HTL) response. Modified erythropoietin (EPO) construct insects are useful for the construction of baculovirus and eukaryotic expression vectors. The present sequence represents human erythropoietin.

SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 8; Length 193;
 Best Local Similarity 100.0%; Pred No. 2, 88-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRLVLEAKAENTTGCAREHCSLNENITWPDTKTYAWKRMVEGQA 60
 DB 28 APPRLICDSRLVLEAKAENTTGCAREHCSLNENITWPDTKTYAWKRMVEGQA 87

QY 61 VEWQGALLSEAVURGQALLVNSQPWEPLQLHVDAVSLRSLTLLAIGAKEYAIS 120
 DB 88 VEWQGALLSEAVURGQALLVNSQPWEPLQLHVDAVSLRSLTLLAIGAKEYAIS 147

QY 121 PPDAAASAPLRTITADFRKLFPRVYSNPLRGKLKUYGAEACTGD 165
 DB 148 PPDAAASAPLRTITADFRKLFPRVYSNPLRGKLKUYGAEACTGD 192

RESULT 82

ID ADT07730 standard; protein; 193 AA.
 XX

AC ADT07730;

DT 13-JAN-2005 (first entry)

XX Human wild-type erythropoietin protein.

XX

KW Erythropoietin; EPO; reduced immunogenicity; reduced immunity;

KW major histocompatibility complex class II; MHC;

KW helper T lymphocyte response; HTL; fungal disease; viral disease;

KW bacterial disease; parasitic disease; cancer; autoimmune disease;

KW alloraft rejection; allergy; Lyme disease; ulcerative colitis;

KW transplantation; haemophilia; osteoporosis; metabolic disease;

KW food hypersensitivity; cytostatic; immunosuppressive; antiinflammatory;

KW human.

OS Homo sapiens.

XX

Key Location/Qualifiers

FP Peptide 1..27

FP Protein /label= Signal_peptide

FP Protein /label= Mature_EPO_protein

XX WO2004089973-A2.

XX 21-OCT-2004.

XX PR 02-APR-2004; 2004WO-US010353.

XX PR 02-APR-2003; 2003US-0459939P.

PA (EPIM-) BIIMMUNE INC.

XX Tangri S, Mothe B, Sette A, Southwood S, Briggs K, Chestnut RW;

DR WPI; 2004-748719/73.

XX New isolated or purified modified erythropoietin construct useful for treatment of anemia comprises a sequence selected from 5 sequences each containing 193 amino acids as given in specification, or truncated modified erythropoietin.

PS Example 1; SEQ ID NO 9; 223pp; English.

XX

CC The invention relates to isolated or purified modified erythropoietin constructs.

CC (EPO) constructs (MEC), and truncated modified erythropoietin constructs.

CC These constructs are peptides, polypeptides, proteins or antibodies

CC having reduced immunogenicity as compared to the naturally occurring

CC form. Also disclosed is a method of producing such peptides. The reduced

CC immunity is as a result of reduced binding to major histocompatibility

CC complex (MHC) class II molecules. The peptides of the invention are

CC useful for antagonising the erythropoietin (EPO) receptor or treating

CC diseases or conditions associated with over-activation of the EPO

CC receptor. The invention is useful for producing a peptide, polypeptide,

CC protein and antibody having reduced immunogenicity, which is useful in

CC the treatment and diagnosis of disease, conditions and disorders. It is

CC also useful for reducing the helper T lymphocyte (HTL) response against a

CC candidate protein. The peptides, polypeptides, proteins and antibodies

CC are useful for the treatment of pathological states (such as fungal,

CC viral, bacterial and parasitic diseases, cancer (such as breast cancer,

CC non-Hodgkin's lymphoma), autoimmune diseases (such as rheumatoid

CC arthritis, multiple sclerosis, myasthenia gravis), allograft rejection,

CC allergies (e.g. pollen allergies), Lyme disease, hepatitis B and C, ICMV,

CC post-streptococcal endocarditis or glomerulonephritis, ulcerative

CC colitis, Crohn's disease, psoriasis, chronic renal failure, asthma,

CC transplantation, haemophilia, Paget's disease, osteoporosis, chronic

CC granulomatous disease, genital warts, diabetes, defective tissue growth,

CC metabolic disease and food hypersensitivities). The peptides, polypeptides,

CC proteins and antibodies are modified so as to have reduced

CC immunogenicity as a result of reduced binding to MHC class II against

CC various DR and DQ molecules and the subsequent reduced helper T

CC lymphocyte (HTL) response. Modified erythropoietin (EPO) construct

CC inserts are useful for the construction of bacterial and eukaryotic

CC expression vectors. The present sequence represents human wild-type

CC erythropoietin.

XX

SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 8; Length 193;

Best Local Similarity 100.0%; P-Value 2.8e-86;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 APPRLICOSRVEYLEENKEAENITTCGCAEHCSLNENITTPDKVNLYAKWKEVMEVGQA 60

Db 28 APPRLICDSRVEYLEENKEAENITTCGCAEHCSLNENITTPDKVNLYAKWKEVMEVGQA 87

Oy 61 VEWVOGLALISSEAVTRGQALLVNSQPMLQLHDKAVSGLSLTTLRAGQAEKLS 120

Db 88 VEWVOGLALISSEAVTRGQALLVNSQPMLQLHDKAVSGLSLTTLRAGQAEKLS 147

Oy 121 PPDAASAAARLTITADTRKLFRYVSNPFLRGKLUYTSACRTGD 165

Db 148 PPDAASAAARLTITADTRKLFRYVSNPFLRGKLUYTGACRTGD 192

RESULT 83

ID ADT9640 standard; protein; 193 AA.

ID ADT9640 standard; protein; 193 AA.

AC ADT9640;

XX

DR 13-JAN-2005 (first entry)

DE Erythropoietin (EPO) receptor seqid 10.

XX respiratory; cardiotropic; vasoconstrictive; CNS; antibacterial;

XX nootropic; immunosuppressive; antiallergic; cytoprotective; osteoprotective;

XX anti-parkinsonian; neuroprotective; antiarrhythmic; antirheumatic;

XX nephrotoxic; muscle activity; thrombolytic; antidiabetic;

XX tissue protective activity; tissue protective cytokine receptor complex;

XX nervous system disorder; hypoxia; ischaemia; epilepsy;

XX chronic seizure disorder; neurotoxin poisoning; septic shock;

XX anaphylactic shock; neuropsychologic disorder; senile dementia;

XX Alzheimer's disease; Parkinson's disease; dementia; multiple sclerosis;

XX Creutzfeldt-Jakob disease; Huntington's disease; inflammatory disease;

KW chronic bronchitis; rheumatoid arthritis; glomerulonephritis;

KW encephalitis; meningitis; polymyositis; ophthalmic disease; angitis;

KW retinal ischaemia; cardiovascular disease; myocardial infarction;

KW myocarditis; cardiopulmonary disease; asthma; pulmonary thrombosis;

KW respiratory disease; kidney disease; urinary disease;

KW reproductive disease; myasthenia gravis; diabetes; autoimmune disease;

KW bone disease; osteopenia; Paget's disease; gastrointestinal disease;

KW endocrine abnormality; metabolic abnormality; complexity; ligand; human;

KW erythropoietin; EPO.

OS Homo sapiens.

XX

PN US2004214236-A1.

XX

PR 25-APR-2003; 2003US-046581P.

XX

PR 30-SEP-2003; 2003US-00676694.

XX

PR 28-OCT-2004.

XX

PT PT Identifying compound modulating tissue protective activity, by contacting test compound with tissue protective cytokine receptor complex, measuring activity level of complex.

XX

PS Disclosure; SEQ ID NO 10; 148PP; English.

XX

CC The invention describes a method of identifying (M1) a compound that modulates tissue protective activity, by contacting test compound with tissue protective cytokine receptor complex (I), measuring the level of activity of (I), identifying test compound that increases/decreases level of activity of (I) as compared to level of activity of (II) measured in absence of the test compound, and assaying identified test compound for tissue protective activity. (M1) is useful for identifying a compound that modulates a tissue protective activity. Also described is a method (M2) for identifying a compound that binds to (I) and a method (M3) for identifying a compound that modulates the binding of a tissue protective cytokine receptor complex ligand to (I), or compound that modulates the interaction between (I) and tissue protective cytokine receptor complex ligand. The compounds identified using (M1)-(M3) are useful for treating various conditions of the central and peripheral nervous systems (e.g., hypoxia, and/or ischaemia, epilepsy, chronic seizure disorders, neurotoxin poisoning, septic shock, anaphylactic shock), neuropsychologic disorders (senile dementia, Alzheimer's disease, Parkinson's disease, dermatoxanthomatosis, multiple sclerosis, Creutzfeldt-Jakob disease, Huntington's disease), inflammatory diseases (e.g., chronic bronchitis, rheumatoid arthritis, glomerulonephritis, encephalitis, meningitis, polymyositis), ophthalmic diseases (e.g., angiitis, retinal ischaemia), cardiovascular diseases (e.g., myocardial infarction, myocarditis), cardiopulmonary diseases (e.g., asthma, pulmonary thrombosis), respiratory diseases, kidney, urinary, and reproductive diseases (e.g., myasthenia gravis, diabetes, autoimmune disease), bone diseases (e.g., osteopenia, Paget's disease), gastrointestinal diseases and endocrine and metabolic abnormalities. (M1) enables identification of compounds that have a tissue protective activity using a heteromultimer receptor complex that mediates the tissue protective activities. This is

CC	the amino acid sequence of human tissue protective cytokine receptor	PA	(PEDE/)	PEDERSEN L.O.
CC	complex ligand erythropoietin (EPO).	XX	"	"
XX		PT	Brines M,	Cerami A,
SQ	Sequence 193 AA;	PI	Fioraliso P,	Fratelli M,
		DR	Nielsen M,	Leist M;
Query Match	100.0%; Score 846; DB 8; Length 193;	XX	WPI; 2004-765699/75.	
Best Local Similarity	100.0%; Pred. No. 2.8e-86;	PT		
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	XX		
QY	1 APRILICDSRVLERYLLEAKENITGCAEHCSLENINITYPDTKNFYAWKRMVQQA 60	CC	Identifying compound modulating tissue protective activity, by contacting	
Db	28 APRILICDSRVLERYLLEAKENITGCAEHCSLENINITYPDTKNFYAWKRMVQQA 87	CC	test compound with tissue protective cytokine receptor complex, measuring	
QY	61 VEWQGLALLSEAVLKGQALLVNSQWEPLQLHVDAVGLSLTLLRAKGKAIS 120	CC	activity level or complex, identifying test compound modulating activity	
Db	88 VEWQGLALLSEAVLKGQALLVNSQWEPLQLHVDAVGLSLTLLRAKGKAIS 147	CC	level of complex.	
QY	121 PDDAASAPLRTITADPFLKLFYVSNFLRGKLYTGACRTGD 165	CC	Disclosure: SEQ ID NO 22; 148pp; English.	
Db	148 PDDAASAPLRTITADPFLKLFYVSNFLRGKLYTGACRTGD 192	CC	The invention describes a method of identifying (M1) a compound that	
RESULT 84		CC	modulates tissue protective activity, by contacting test compound with	
ID ADT9562		CC	tissue protective cytokine receptor complex (II), measuring the level of	
ID ADT99652 standard; protein; 193 AA.		CC	activity of (I), identifying test compound that increases/decreases level	
AC ADT98652;		CC	of activity of (II) as compared to level of activity of (I) measured in	
XX		CC	absence of the test compound, and assaying identified test compound for	
DT 13-JAN-2005 (first entry)		CC	tissue protective activity. (M1) is useful for identifying a compound	
XX		CC	that modulates a tissue protective activity. Also described is a method	
DE Brythropoietin (EPO) receptor mutant seqid 22.		CC	(M2) useful for identifying a compound that binds to (I) and a method	
XX		CC	(M3) for identifying a compound that modulates the binding of a tissue	
XX		CC	protective cytokine receptor complex ligand to (I), or compound that	
XX		CC	modulates the interaction between (I) and tissue protective cytokine	
XX		CC	receptor complex ligand. The compounds identified using (M1)-(M3) are	
XX		CC	useful for treating various conditions of the central and peripheral	
XX		CC	nervous systems (e.g., hypoxia, ischemia, epilepsy, chronic	
XX		CC	seizure disorders, neurotoxin poisoning, septic shock, anaphylactic	
XX		CC	shock), neuropsychologic disorders (senile dementia, Alzheimer's disease,	
XX		CC	Parkinson's disease, dementia, multiple sclerosis, Creutzfeldt-Jakob	
XX		CC	disease, Huntington's disease), inflammatory diseases (e.g. chronic	
XX		CC	bronchitis, rheumatoid arthritis, glomerulonephritis, encephalitis,	
XX		CC	meningitis, polyradiculitis), ophthalmic diseases (e.g., retinal	
XX		CC	ischemia), cardiovascular diseases (e.g., myocardial infarction,	
XX		CC	myocarditis), cardiopulmonary diseases (e.g., asthma, pulmonary	
XX		CC	thrombosis), respiratory diseases, kidney, urinary, and reproductive	
XX		CC	diseases (e.g., myasthenia gravis, diabetes, autoimmune diseases), bone	
XX		CC	diseases (e.g., osteopenia, Paget's disease), gastrointestinal diseases	
XX		CC	and endocrine and metabolic abnormalities. (M1) enables identification of	
XX		CC	compounds that have a tissue protective activity using a heteromultimer	
XX		CC	receptor complex that mediates the tissue protective activities. This is	
XX		CC	the amino acid sequence of a human tissue protective cytokine receptor	
OS Sequence 193 AA;		CC	complex ligand erythropoietin (EPO) mutant.	
OS		XX		
OS Synthetic.		XX		
XX		XX		
US2004214336-A1.		XX		
XX		XX		
PD 28-OCT-2004.		XX		
XX		XX		
PP 30-SEP-2003; 2003US-00676894.		XX		
PR 25-APR-2003; 2003US-0465891P.		XX		
XX		XX		
PA (BRIN/ BRINES M.		XX		
PA (CERA/ CERAMI A.		XX		
PA (GHEZ/ GHIZZI P.		XX		
PA (FIOR/ FIORDALISO P.		XX		
PA (FRAT/ FRATELLI M.		XX		
PA (LEIS/ LEIST M.		XX		
PA (NIEL/ NIELSEN M.		XX		
PA (SAGE/ SAGER T.		XX		
PA (GERW/ GERWIEN J.		XX		
RESULT 85		XX		
ADT99742		XX		
ID ADT99742 standard; protein; 193 AA.		XX		
AC ADT99742;		XX		
XX		XX		
DT 13-JAN-2005 (first entry)		XX		

DE Brythropoietin (EPO) receptor mutant seqid 112.
 XX respiratory; cardiotropic; anticonvulsant; CNS; antibacterial;
 KW nootropic; immunosuppressive; antiallergic; cyrostatic; osteopathic;
 KW antiparkinsonian; neuroprotective; antiarthritic; antirheumatic;
 KW nephropathic; muscular; thrombolytic; antidiabetic;
 KW tissue protective activity; tissue protective cytokine receptor complex;
 KW nervous system disorder; hypoxia; ischaemia; epilepsy;
 KW chronic seizure; neurotoxin poisoning; septic shock;
 KW anaphylactic shock; neuropsychologic disorder; senile dementia;
 KW Alzheimer's disease; Parkinson's disease; dementia; multiple sclerosis;
 KW Creutzfeldt-Jakob disease; Huntington's disease; inflammatory disease;
 KW chronic bronchitis; rheumatoïd arthritis; glomerulonephritis; chronic
 KW encephalitis; meningitis; polymyositis; opthalmalic disease; encephalitis;
 KW retinal ischaemia; cardiovascular disease; myocardial infarction;
 KW myocarditis; cardiopulmonary disease; asthma; pulmonary thrombosis;
 KW respiratory disease; kidney disease; urinary disease;
 KW bone disease; osteopenia; Paget's disease; gastrointestinal disease;
 KW endocrine abnormality; metabolic abnormality;
 KW tissue protective cytokine receptor complex ligand; human;
 XX OS Homo sapiens.
 XX Synthentic.
 XX US2004214236-A1.
 XX PD 28-OCT-2004.
 XX PR 30-SEP-2003; 2003US-00676694.
 XX PR 25-APR-2003; 2003US-0465891P.
 XX PA (BRIN/) BRINES M.
 PA (CERA/) CERAMI A.
 PA (GHEZ/) GHEZZI P.
 PA (FIOR/) FIORDALISO P.
 PA (FRAT/) FRATELLI M.
 PA (LEIS/) LEIST M.
 PA (NIEL/) NIELSEN M.
 PA (SAGE/) SAGER T.
 PA (GERW/) GERWIEN J.
 PA (PDE/) PEDERSEN L O.
 XX PI Brines M., Cerami A., Ghezzi P., Fiordaliso P., Fratelli M., Leist M.;
 PT Nielsen M., Sager T., Gerwiens J., Pedersen LO;
 XX DR WPI; 2004-765609/75.
 XX PS Disclosure; SEQ ID NO 112; 148pp; English.

The invention describes a method of identifying (M1) a compound that modulates tissue protective activity, by contacting test compound with tissue protective cytokine receptor complex (I), measuring the level of activity of (I), identifying test compound that increases/decreases level of activity of (I) as compared to level of activity of (I) measured in absence of the test compound, and assaying identified test compound for tissue protective activity. (M1) is useful for identifying a compound that modulates a tissue protective activity. Also described is a method (M2) for identifying a compound that binds to (I) and a method (M3) for identifying a compound that modulates the binding of a tissue protective cytokine receptor complex ligand to (I), or compound that modulates the interaction between (I) and tissue protective cytokine receptor complex ligand. The compounds identified using (M1)-(M3) are useful for treating various conditions of the central and peripheral nervous systems (e.g., hypoxia, and/or ischaemia, epilepsy, chronic seizure disorders, neurotoxin poisoning, septic shock, anaphylactic

CC shock), neuropsychologic disorders (senile dementia, Alzheimer's disease, Parkinson's disease, dermititis, multiple sclerosis, Cratzfeldt-Jakob disease, Huntington's disease), inflammatory diseases (e.g., chronic bronchitis, rheumatoid arthritis, glomerulonephritis, encephalitis, meningoencephalitis, polymyositis), ophthalmalic diseases (e.g., arglitis, retinal ischaemia), cardiovascular diseases (e.g., myocardial infarction, myocarditis), cardiopulmonary diseases (e.g., asthma, pulmonary thrombosis), respiratory diseases, kidney, urinary, and reproductive diseases (e.g., myasthenia gravis, diabetes, autoimmune diseases), bone diseases (e.g., osteopenia, Paget's disease), gastrointestinal diseases and endocrine and metabolic abnormalities. (M1) enables identification of compounds that have a tissue protective activity using a heteromultimer receptor complex that mediates the tissue protective activities. This is the amino acid sequence of a human tissue protective cytokine receptor complex ligand erythropoietin (EPO) mutant.

XX SQ Sequence 193 AA;

Query Match 100.0%; score 846; DB 8; Length 193;
 Best Local Similarity 100.0%; Pred. 2. 8e-6; Mismatches 0; Indels 0; Gaps 0; Matches 165; Pred. 1. 0; Gaps 0; Indels 0; Mismatches 0.

Qy 1 APPRLIDDSRVLYVYLEAKRBNITTCACBHSCLNNNTIVDTKPNFYAWKGMEMVQQAA 60
 Db 28 APPRLIDDSRVLYVYLEAKRBNITTCACBHSCLNNNTIVDTKPNFYAWKGMEMVQQAA 87

Qy 61 VEVWQGLLSEAVLRLGQALYNSSQMPWEPLOHVADAVSGRSLSLTTRALGAQKRAIS 120
 Db 88 VEVWQGLLSEAVLRLGQALYNSSQMPWEPLOHVADAVSGRSLSLTTRALGAQKRAIS 147

Qy 121 PPDASASAPLRTTADTPRKLFRVYSNPLRGKLUKYGEACRGD 165
 Db 148 PPDASASAPLRTTADTPRKLFRVYSNPLRGKLUKYGEACRGD 192

RESULT 86

ID AEB92238 standard; protein; 193 AA.

XX AC AEB92238;

XX DT 06-OCT-2005 (first entry)

DB Erythropoietin, SEQ ID 10.

XX Antianemic; Gene therapy; anemia; erythropoietin.

OS Homo sapiens.

XX PN US2005158822-A1.

XX PD 21-JUL-2005.

XX PF 20-JAN-2004; 2004US-00759031.

PR 20-JAN-2004; 2004US-00759031.

XX (INSI-) INSIGHT BIOPHARMACEUTICALS LTD.

XX Pecker I;

XX WPI; 2005-589511/60.

DR N_PSDB; AEB92236, AEB92237, AEB92239, AEB92240.

DR RFPSEQ; NP_000790.

XX New chimeric Polynucleotide comprises a nucleic acid encoding an erythropoietin (EPO) polypeptide attached to a 5'-UTR sequence, useful for producing high levels of EPO in mammalian cells for treating disorders, e.g. anemia.

XX Claim 2; SEQ ID NO 10; 24pp; English.

XX The present invention relates to a novel chimeric polynucleotide (I),

CC which comprises a nucleic acid sequence encoding an erythropoietin (EPO) protein (AEB92238) attached to a 5'-UTR sequence (AEB9234 or AEB9235).
 CC The 5'-UTR sequences improve the translational efficiency of fused EPO coding sequences in eukaryotic cells. In addition, to further improve the CC translation activity of (I), the GC content of the sequence can be reduced. This was illustrated by AEB9237, where the GCG triplet encoding CC the Glycine residue at position 2 of EPO protein, was mutated to GGA, via CC a guanine to adenine substitution. (I) is useful for producing high levels of EPO in mammalian cells and can be used to treat disorders, such as CC which are associated with, or lead to, abnormal EPO production, such as CC anemia.
 XX SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 9; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 OY 1 APPRLICDSRVLEERYLLEAKERENITGCAEHCSLNENITVPDTKIVFYAWKRMEVQQA 60
 DB 28 APPRLICDSRVLEERYLLEAKERENITGCAEHCSLNENITVPDTKIVFYAWKRMEVQQA 87
 OY 61 VEWVGALLSEAVLRGQALVNNSOPWEPLOHLVDAVKGAVSGLSLTLLRAIGAKEAIIS 120
 DB 88 VEWVGALLSEAVLRGQALVNNSOPWEPLOHLVDAVKGAVSGLSLTLLRAIGAKEAIIS 147
 OY 121 PPDASASAPLRTITADTFRKLYVSNPLRGKLYTGACRTGD 165
 DB 148 PPDASASAPLRTITADTFRKLYVSNPLRGKLYTGACRTGD 192

RESULT 87

ABC05272 standard; protein: 193 AA.

AC ABC05272;

XX DT 06-OCT-2005 (first entry)

DB Human precursor erythropoietin polypeptide.

KW Hormone; erythropoietin; anemia; renal failure; cerebral ischemia; brain injury; spinal cord injury; retinopathy; Alzheimer's disease; Parkinsons disease; Huntingtons chorea; motor neurone disease; sickle cell anemia; beta thalassemia; cystic fibrosis; pregnancy disorder; menstruation disorder; aging; antianemic; nephroprotective; cerebroprotective; vasotrophic; neuroprotective; vulnerability; ophthalmological; nootropic; antiparkinsonian; anticonvulsant; dermatological; CNS-Gen.; muscular-gen.; respiratory-gen.; gynecological; antischiking; CNS-Gen.; muscular-gen.; respiratory-gen.; gynecological; dermatological.

OS Homo sapiens.

XX PN WO2005065239-A2.

XX PD 21-JUL-2005.

PP 23-DEC-2004; 2004WO-US043081.

PR 31-DEC-2003; 2003US-0533617P.

XX PA (CENZ) CENTOCOR INC.

XX PI Pool C, Mills J, Cunningham M;

DR WPI; 2005-618232/63.

KW Erythropoietic conjugate for treating anemia, retinal disease, Alzheimer's disease, Parkinson's disease, Huntington's disease, has N-terminal free thiols, and capable of causing bone marrow cells to increase production of red blood cells.

PS Disclosure; SEQ ID NO 14; 57pp; English.

XX

The invention relates to an erythropoietic (EPO) conjugate capable of causing bone marrow cells to increase production of red blood cells. The EPO conjugate contains recombinant/non-recombinant mammalian erythropoietin in which a cysteine residue having a free alpha amine has been added by recombinant, enzymatic or chemical means, to provide a reactive free thiol that does not interfere with protein folding, secretion or bioactivity and thiol may be derived, thus increasing the circulating half-life or improving the biological activity of the erythropoietic protein. The invention also relates to a method of preparing a therapeutic protein conjugate having a polymer conjugated to the N-terminal cysteine of the therapeutic protein, where the thiol of the cysteine residue participates in formation of a covalent bond of the conjugate, involving obtaining a nucleic acid sequence for the therapeutic protein in a cell and obtaining a nucleic acid sequence for the signal sequence, directing the formation of a construct by engineering of the signal sequence to the therapeutic protein sequence with the codon TGT interposed between them so that the signal sequence is upstream of the TGT causing the construct to be expressed in the cell, recovering the polypeptide coded for by the construct and conjugating the polypeptide at the N-terminal cysteine to a polymer, and preparing the EPO conjugate by contacting a cys-EPO moiety having a cysteine residue at the N-terminus with a preconstructed hydrophilic polymer-organic moiety. The EPO conjugate is useful for treating anemia, renal failure, cerebral ischemia, brain injury, spinal cord injury, retinal disease, Alzheimer's disease, Parkinson's disease, Huntington's disease, amyotrophic lateral sclerosis, sickle cell disease, beta thalassemia, cystic fibrosis, pregnancy disorders, menstrual disorders and aging. This sequence represents a human precursor erythropoietin polypeptide used in the scope of the invention.

XX SQ Sequence 193 AA;

Query Match 100.0%; Score 846; DB 9; Length 193;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 APPRLICDSRVLEERYLLEAKERENITGCAEHCSLNENITVPDTKIVFYAWKRMEVQQA 60

DB 28 APPRLICDSRVLEERYLLEAKERENITGCAEHCSLNENITVPDTKIVFYAWKRMEVQQA 87

OY 61 VEWVGALLSEAVLRGQALVNNSOPWEPLOHLVDAVKGAVSGLSLTLLRAIGAKEAIIS 120

DB 88 VEWVGALLSEAVLRGQALVNNSOPWEPLOHLVDAVKGAVSGLSLTLLRAIGAKEAIIS 147

OY 121 PPDASASAPLRTITADTFRKLYVSNPLRGKLYTGACRTGD 165

DB 148 PPDASASAPLRTITADTFRKLYVSNPLRGKLYTGACRTGD 192

RESULT 88

ABC05259 standard; protein: 193 AA.

AC ABC05259;

XX DT 06-OCT-2005 (first entry)

DB Human erythropoietin polypeptide.

XX Hormone; erythropoietin; anemia; renal failure; cerebral ischemia;

KW brain injury; spinal cord injury; retinopathy; Alzheimer's disease;

KW Parkinsons disease; Huntingtons chorea; motor neurone disease;

KW sickle cell anemia; beta thalassemia; cystic fibrosis; pregnancy disorder; menstruation disorder; aging; antianemic; nephroprotective; cerebroprotective; vasotrophic; neuroprotective; vulnerability; ophthalmological; nootropic; antiparkinsonian; anticonvulsant; dermatological; CNS-Gen.; muscular-gen.; respiratory-gen.; gynecological;

PS Disclosure; SEQ ID NO 14; 57pp; English.

OS Homo sapiens.

XX

RESULT 90
 AAW62048
 ID AAW62048 standard; protein; 194 AA.
 XX
 AC AAW62048;
 XX DT 10-SEP-1998 (first entry)
 DE Human erythropoietin clone 7.2.
 XX KW Human; erythropoietin; EPO; Chinese hamster ovary cell; CHO; strain;
 KW medicine; biological research.
 XX OS Homo sapiens.
 XX FH Key Peptide
 PT 1.:27
 PT /label= signal
 FT Protein
 FT 28.:194
 PT /label= erythropoietin
 PN RU2089611-C1.
 PD 10-SEP-1997.
 XX PP 13-JUL-1995; 95RU-00111858.
 XX PR 13-JUL-1995; 95RU-00111858.
 XX PA (MED BIOTECHN RES PRODN CENTRE.
 XX PT Zelenin MG, Kamerova IA, Kolobkov SI;
 XX DR WPI; 1998-205757/18.
 DR N-PSDB; AAV37951.
 XX PT New strain of cultivated cells of Chinese hamster - acts as producer of
 PT human erythropoietin which can be used in medicine and in biological
 PT research.
 XX PS Disclosure; Col 15-22; 13pp; English.
 CC The present sequence represents human erythropoietin clone 7.2 from the
 CC present invention. The present invention describes a new CHO strain of
 CC cultivated cells of Chinese hamster VSK (P) 637 D, which produces human
 CC erythropoietin. The new strain is used as a new strain-producer of human
 CC erythropoietin, which can be used in medical therapy and research, and
 CC also in biological research. The use of the strain reduces the cost of
 CC production of human erythropoietin owing to increased productivity of the
 CC strain.
 SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 2; Length 194;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; DB Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APRPLICDSRVLERYLILBAKEAENITGCAEHCISLNENITVPTDKVNFYAWRMEVGQA 60
 Db 29 APRPLICDSRVLERYLILBAKEAENITGCAEHCISLNENITVPTDKVNFYAWRMEVGQA 88
 Qy 61 VEWQGLALLSEAVLRGOALLVNSSQWEPLOLHVDAVSGRLSLTLLRAIGAKEAI 120
 Db 89 VEWQGLALLSEAVLRGOALLVNSSQWEPLOLHVDAVSGRLSLTLLRAIGAKEAI 148
 Qy 121 PPDASAPLRTITADTFKLFRVYSIFLRGKLYTGBCRTGD 165
 Db 149 PPDASAPLRTITADTFKLFRVYSIFLRGKLYTGBCRTGD 193

RESULT 92
 ADL06826
 ID ADL06826 standard; protein; 194 AA.
 XX AC ADL06826;
 XX DT 03-JUN-2004 (first entry)
 XX OS Homo sapiens.

DB19855489-A1.
 XX PD 17-AUG-2000.
 XX PF 01-DEC-1998; 98DE-01055489.
 XX PR 01-DEC-1998; 98DE-01055489.
 XX PA (GROZ/) GROZA I.
 XX DR WPI; 2000-566040/53.
 DR N-PSDB; AAV71992.

XX PT New nucleic acid molecule comprising simian virus 40 regulatory sequences
 PT and antibiotic resistance gene, useful for expressing erythropoietin in
 PR mammalian cells for treating anemia.
 XX PS Claim 1; Fig 5; 18pp; German.

CC This invention describes a novel nucleic acid molecule (I) encoding an
 CC erythropoietin (EPO) polypeptide (II), transcriptional and translational
 CC regulatory sequences from simian virus 40 (SV40), including the SV40
 CC early promoter and a sequence encoding resistance to an antibiotic. The
 CC product of the invention has anti-anemic activity. EPO regulates
 CC proliferation and differentiation of late erythrocyte precursor cells.
 CC (I) is used for the recombinant production of human EPO in mammalian
 CC cells. EPO is used in replacement therapy, to treat anemia. Cells
 CC transformed with (I) produce EPO at a high level (e.g. 150-180
 CC international units/ml) which is stable under non-selection conditions.
 CC The plasmid copy number in the cells can be increased without using the
 CC expensive and highly cytostatic agent methotrexate. This sequence
 CC represents the human erythropoietin protein which is described in the
 XX methods of the invention

SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 3; Length 194;
 Best Local Similarity 100.0%; Pred. No. 2.8e-86; DB Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APRPLICDSRVLERYLILBAKEAENITGCAEHCISLNENITVPTDKVNFYAWRMEVGQA 60
 Db 29 APRPLICDSRVLERYLILBAKEAENITGCAEHCISLNENITVPTDKVNFYAWRMEVGQA 88
 Qy 61 VEWQGLALLSEAVLRGOALLVNSSQWEPLOLHVDAVSGRLSLTLLRAIGAKEAI 120
 Db 89 VEWQGLALLSEAVLRGOALLVNSSQWEPLOLHVDAVSGRLSLTLLRAIGAKEAI 148
 Qy 121 PPDASAPLRTITADTFKLFRVYSIFLRGKLYTGBCRTGD 165
 Db 149 PPDASAPLRTITADTFKLFRVYSIFLRGKLYTGBCRTGD 193

RESULT 91
 AAB10654
 ID AAB10654 standard; protein; 194 AA.

RESULT 91
 AAB10654
 ID AAB10654 standard; protein; 194 AA.

DE Human 165 residue erythropoietin analogue #45.

XX Human; erythropoietin; EPO; iron distribution disturbance; diabetes;

KW non-insulin dependent diabetes; type 2 diabetes; reticulocyte production;

KW red blood cell production; glycosylation site; analogue; antidiabetic;

KX mutant; mutein.

OS Homo sapiens.

Synthetic.

PN WO2004019972-A1.

PD 11-MAR-2004.

XX 20-AUG-2003; 2003WO-EP009194.

XX 29-AUG-2002; 2002EP-00019100.

PR XX (HOFF) HOFFMANN LA ROCHE & CO AG F.

PA XX PT Lehmann P., Roediger R., Walter-Matsui R;

XX DR XX

PT XX

PS XX

DR XX

PT Use of erythropoietin protein in manufacture of medicament for treating disturbances of iron distribution in diabetes.

PS Disclosure; Page, 31pp; English.

The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in diabetes. The

activation or an erythropoietin analogue such as darbepoietin alpha. The

erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation.

Patients with diabetes have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall

concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effects of iron accumulation in certain organs, leading to organ damage and destruction, and/or experience effects similar to anaemia due to iron usage in blood

cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticulocytes and red blood cells, and this has been found to have a beneficial effect on iron distribution disturbances in diabetes e.g., non-insulin dependent (type 2) diabetes. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron distribution in diabetes. Sequences ADU06807-ADU06831 represent analogues of the 166 amino acid human erythropoietin which contain additional or altered glycosylation sites.

Note: The present sequence is not shown in the specification, but is derived from the wild-type 166 residue human EPO (ADU06781) and the information given on page 6.

XX SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 8; Length 194;

Best Local Similarity 100.0%; Pred. No. 2. 8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

QY 61 VEVVOGLALLSEAVRQQLVNSQWPWLQLHVDKAVSGLRLTILRAQGQEKAIS 120

CC SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 8; Length 194;

Best Local Similarity 100.0%; Pred. No. 2. 8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

61 VEVVOGLALLSEAVRQQLVNSQWPWLQLHVDKAVSGLRLTILRAQGQEKAIS 120

CC SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 8; Length 194;

Best Local Similarity 100.0%; Pred. No. 2. 8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

61 VEVVOGLALLSEAVRQQLVNSQWPWLQLHVDKAVSGLRLTILRAQGQEKAIS 120

RESULT 93

ID AD059461

ID AD059461 standard; protein; 194 AA.

AC AD059461;

DT 26-AUG-2004 (first entry)

XX Human 165 residue erythropoietin analogue #45.

KW Human; erythropoietin; EPO; iron distribution disturbance; heart disease; heart insufficiency; coronary heart disease; atherosclerosis; acute coronary syndrome; heart failure; congestive heart failure; reticulocyte production; red blood cell production; cardiac; antiarteriosclerotic; glycosylation site; analogue; mutant; mutein.

OS Homo sapiens.

Synthetic.

PN WO2004047858-A1.

XX PR XX

XX PD 10-JUN-2004.

XX PA XX

XX PR 17-NOV-2003; 2003WO-EP012822.

XX PR 22-NOV-2002; 2002EP-00026342.

XX PA XX

XX PR (HOFF) HOFFMANN LA ROCHE & CO AG F.

XX PI Lehmann P., Roediger R., Walter-Matsui R;

XX DR XX

XX PS XX

XX Use of erythropoietin protein in the manufacture of medicament for treating disturbances of iron distribution in heart diseases e.g. heart insufficiency.

PT WPI; 2004-450212/42.

XX Disclosure; Page, 31pp; English.

The invention relates to the use of an erythropoietin (EPO) protein for the treatment of disturbances of iron distribution in heart diseases. The

activation or an erythropoietin analogue such as darbepoietin alpha. The

erythropoietin protein used in the method may also be modified by the addition of 1-6 glycosylation sites, or by pegylation.

Patients with heart diseases have been found to have a high probability of being affected by disturbances of iron distribution. In such patients, the overall

concentration of iron in the body is normal (compared with conditions such as anaemia), but the individual may suffer the effect of iron accumulation in certain organs, leading to organ damage and destruction,

and/or experience effects similar to anaemia due to iron usage in blood

cell formation being impaired. Erythropoietin causes bone marrow cells to increase production of reticulocytes and red blood cells, and this has

been found to have a beneficial effect on iron distribution disturbances in heart diseases e.g., heart insufficiency, coronary heart disease, atherosclerosis, acute coronary syndrome, heart failure and congestive

heart failure. Erythropoietin proteins may therefore be used to manufacture a medicament for the treatment of disturbances of iron

distribution in heart diseases. Sequences AD059442-AD059466 represent

analogues of the 166 amino acid human erythropoietin which contain additional or altered glycosylation sites. Note: The present sequence is not shown in the specification, but is derived from the wild-type 166

residue human EPO (AD059416) and the information given on page 6.

XX SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 8; Length 194;

Best Local Similarity 100.0%; Pred. No. 2. 8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

61 VEVVOGLALLSEAVRQQLVNSQWPWLQLHVDKAVSGLRLTILRAQGQEKAIS 120

CC SQ Sequence 194 AA;

Query Match 100.0%; Score 846; DB 8; Length 194;

Best Local Similarity 100.0%; Pred. No. 2. 8e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLETVLLEAKEAENITTCGCAEHCSLNENTIVDPDKYFVAKWKGMEVQQA 60

61 VEVVOGLALLSEAVRQQLVNSQWPWLQLHVDKAVSGLRLTILRAQGQEKAIS 120

Db 1 APRLICDSRIVLERYLLEAKEAENITGCAEHCSLENNTVPTDKINPYAKRMEVQQA 60
Qy 61 VEVMOGLALLSERAVLKGALLNSQWEPLQLHVDAVSGRSLSLTLLRAIGKAIS 120
Db 61 VEVMOGLALLSERAVLKGALLNSQWEPLQLHVDAVSGRSLSLTLLRAIGKAIS 120
Qy 121 PPDASAASAPLRTTADTPRKLFRVVSNFLRGKLUYGEACRTGD 165
Db 121 PPDASAASAPLRTTADTPRKLFRVVSNFLRGKLUYGEACRTGD 165

RESULT 94

ABB77902
ID ABB77902 standard; protein; 196 AA.
XX
AC ABB77902;
XX
DT 07-OCT-2002 (first entry)

DB Amino acid sequence of a modified human erythropoietin (EPO).
KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production; red blood cell production; anaemia; chronic renal failure; acquired immunodeficiency syndrome; AIDS; cancer; bone marrow; committed erythroid progenitor.
KW
OS Synthetic.
OS Homo sapiens.
XX
FH
FT Peptide
FT /note= "secretion signal peptide"
FT Cleavage-site
FT 28..30
FT Protein
FT 31..196
FT /note= "EPO protein"

WO200249673-A2.
XX
PN 27-JUN-2002.
PP 08-DEC-2001; 2001WO-EP014434.
XX
PR 20-DEC-2000; 2000EP-00127891.
PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

PI Burg J, Engel A, Franze R, Hilger B, Schurig HE, Tischer W;
PI Wozny M;
DR WPI; 2002-566640/60.
DR N-PSDB; ABU59290.

XX
PT Novel conjugate of erythropoietin glycoprotein with Polyethylene glycol, useful for treating diseases correlated with anemia in chronic renal failure patients and acquired immunodeficiency syndrome.
PS Disclosure; Fig 4; 40pp; English.

XX
CC The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased circulating half-life and plasma residence time, decreased clearance, increased clinical activity in vivo, improved potency and stability, when compared to unmodified EPO. The EPO conjugate is useful for preparing medicaments for the treatment and prophylaxis of diseases correlated with

RESULT 95

ABB77901
ID ABB77901 standard; protein; 201 AA.
XX
AC ABB77901;
XX
DT 07-OCT-2002 (first entry)

DB Amino acid sequence of a modified human erythropoietin (EPO).
XX
KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production; red blood cell production; anaemia; chronic renal failure; acquired immunodeficiency syndrome; AIDS; cancer; bone marrow; committed erythroid progenitor.
XX
OS Synthetic.
OS Homo sapiens.
XX
FH
FT Peptide
FT 1..27
FT Cleavage-site
FT 28..35
FT Protein
FT 36..201
FT /note= "EPO protein"

WO200249673-A2.
XX
PN 27-JUN-2002.
PP 08-DEC-2001; 2001WO-EP014434.
XX
PR 20-DEC-2000; 2000EP-00127891.
XX
PA (HOFF) HOFFMANN LA ROCHE & CO AG F.

PI Burg J, Engel A, Franze R, Hilger B, Schurig HE, Tischer W;
PI Wozny M;
DR WPI; 2002-566640/60.
DR N-PSDB; ABU59299.

XX
PT Novel conjugate of erythropoietin glycoprotein with polyethylene glycol, useful for treating diseases correlated with anemia in chronic renal failure patients and acquired immunodeficiency syndrome.
PS Disclosure; Fig 3; 40pp; English.

CC anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

SQ Sequence 196 AA;

Query Match	100 %	Score	846	DB	5	Length	196
Best Local Similarity	100.0%	Pred. No.	2.8e-86				
Matches	165	Conservative	0	Mismatches	0	Indels	0
Gaps	0						
Qy	1 APRLICDSRIVLERYLLEAKEAENITGCAEHCSLENNTVPTDKINPYAKRMEVQQA 60						
Db	31 APRLICDSRIVLERYLLEAKEAENITGCAEHCSLENNTVPTDKINPYAKRMEVQQA 90						
Qy	61 VEVMOGLALLSERAVLKGALLNSQWEPLQLHVDAVSGRSLSLTLLRAIGKAIS 120						
Db	91 VEVMOGLALLSERAVLKGALLNSQWEPLQLHVDAVSGRSLSLTLLRAIGKAIS 150						
Qy	121 PPDASAASAPLRTTADTPRKLFRVVSNFLRGKLUYGEACRTGD 165						
Db	151 PPDASAASAPLRTTADTPRKLFRVVSNFLRGKLUYGEACRTGD 195						

The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein is covalently linked to a poly(ethylene glycol) group. The EPO glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased circulating half-life and plasma residence time, decreased clearance, when compared to unmodified EPO. The EPO conjugate is useful for preparing medicaments for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

SQ Sequence 201 AA:

Query Match 100.0%; Score 846; DB 5; Length 201;
 Best Local Similarity 100.0%; Pred. No. 2. 9e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 APPRLICDSRVRLERYLLEAKEAENITTCGAEHCSLNENITVPTDKVNFYAWKRMEVGQQA 60
 Db 36 APPRLICDSRVRLERYLLEAKEAENITTCGAEHCSLNENITVPTDKVNFYAWKRMEVGQQA 95
 QY 61 VEWQGLAILSEAVLVRGQALLVNSQPEPLQHVDKAVSGRSLTTLRALGQKAIS 120
 Db 96 VEWQGLAILSEAVLVRGQALLVNSQPEPLQHVDKAVSGRSLTTLRALGQKAIS 155
 QY 121 PPDASAAAPLRTTADTPKLPRVSNFLRGKLUYTGEACRGD 165
 Db 156 PPDASAAAPLRTTADTPKLPRVSNFLRGKLUYTGEACRGD 200

RESULT 96

ABB77903 ID ABB77903 standard; protein; 201 AA.
 AC ABB77903;
 DT 07-OCT-2002 (first entry)

Amino acid sequence of a modified human erythropoietin (EPO).
 KW Human; erythropoietin; EPO; glycoprotein; reticulocyte production; red blood cell production; anaemia; chronic renal failure; acquired immunodeficiency syndrome; AIDS; cancer; bone marrow; committed erythroid progenitor.

OS Synthetic.

OS Homo sapiens.

FH Location/Qualifiers

FT Peptide 1-.27 /notes= "secretion signal peptide"
 FT Cleavage-site 28-.35 /notes= "proteolytic cleavage site"
 FT Protein 36-.201 /notes= "EPO protein"

PN W0200249673-A2.
 XX 27-JUN-2002.
 PD 08-DEC-2001; 2001WO-EP014434.
 XX PR 20-DEC-2000; 2000EP-00127891.

PA (HOPP) HOFFMANN LA ROCHE & CO AG F.
 XX PI Burg J.; Engel A.; Franz R.; Hilger B.; Schuring HE.; Tischer W.;
 XX PI Wozny M.;
 DR WBI; 2002-566640/60.
 DR N-PSDB; ABL59291.

XX PT Novel conjugate of erythropoietin glycoprotein with polyethylene glycol, useful for treating diseases correlated with anemia in chronic renal failure patients and acquired immunodeficiency syndrome.

XX PS Disclosure; Fig 5; 4app; English.

CC The present sequence represents a modified human erythropoietin (EPO) protein. The EPO was extended at the N-terminal by a proteolytic cleavage site. It was used to produce conjugates of the invention. The specification describes a conjugate comprising an EPO glycoprotein having an N-terminal alpha-amino group, chosen from human EPO (hEPO) or its analogues (where hEPO is modified by addition of 1-6 glycosylation sites or a rearrangement of a glycosylation site). The glycoprotein has in vivo biological activity of causing bone marrow cells to increase production of reticulocytes and red blood cells. The conjugate increased circulating half-life and plasma residence time, decreased clearance, when compared to unmodified EPO. The EPO conjugate is useful for preparing medicaments for the treatment and prophylaxis of diseases correlated with anaemia in chronic renal failure patients (CRF), acquired immunodeficiency syndrome (AIDS) and for treating cancer patients undergoing chemotherapy. It is also useful for treating patients by stimulating the division and differentiation of committed erythroid progenitors in the bone marrow

SQ Sequence 201 AA:

Query Match 100.0%; Score 846; DB 5; Length 201;
 Best Local Similarity 100.0%; Pred. No. 2. 9e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKEAENITTCGAEHCSLNENITVPTDKVNFYAWKRMEVGQQA 60
 Db 36 APPRLICDSRVRLERYLLEAKEAENITTCGAEHCSLNENITVPTDKVNFYAWKRMEVGQQA 95
 QY 61 VEWQGLAILSEAVLVRGQALLVNSQPEPLQHVDKAVSGRSLTTLRALGQKAIS 120
 Db 96 VEWQGLAILSEAVLVRGQALLVNSQPEPLQHVDKAVSGRSLTTLRALGQKAIS 155
 QY 121 PPDASAAAPLRTTADTPKLPRVSNFLRGKLUYTGEACRGD 165
 Db 156 PPDASAAAPLRTTADTPKLPRVSNFLRGKLUYTGEACRGD 200

RESULT 97

ABB05278 ID ABB05278 standard; protein; 201 AA.
 AC ABB05278;
 DT 06-OCT-2005 (first entry)

XX DB Modified human erythropoietin polypeptide.
 XX Hormone; erythropoietin; anemia; renal failure; cerebral ischemia; brain injury; spinal cord injury; retinopathy; Alzheimers disease; Parkinsons disease; Huntington's chorea; motor neurone disease; sickle cell anemia; beta thalassemia; cystic fibrosis; pregnancy disorder; menstruation disorder; aging; antiarrhythmic; nephrotoxic; cerebroprotective; vasotrophic; neuroprotective; pulmonary; ophthalmological; noctropic; antiparkinsonian; anticonvulsant; antiasthmatic; CNS-Gen.; muscular-gen.; respiratory-gen.; gynecological; dermatological; mutein.

XX PA (HOPP) HOFFMANN LA ROCHE & CO AG F.
 XX PI Burg J.; Engel A.; Franz R.; Hilger B.; Schuring HE.; Tischer W.;
 XX PI Wozny M.;
 DR WBI; 2002-566640/60.
 DR N-PSDB; ABL59291.

RESULT 99	Db	100 VEVNQGLALISEAVRQGQALLVNSQPMWPLQHDKAVSGRLSITLRAQKEAIS 159	XX	ID ABB79939 standard; protein; 220 AA.
QY	121 PPDAASAAPLRTTIDTFRKLFRVSNFLRGKLYKTYGACRTGD 165	XX	AC ABB79939;	
Db	160 PPDAASAAPTPITTDFTFKLFRVSNFLRGKLYKTYGACRTGD 204	XX	DE Human erythropoietin-HCG C-terminal peptide fusion protein ECTP;	
ID	AD079063 standard; protein; 209 AA.	DT	KW Human chorionic gonadotropin; HCG; human; erythropoietin; EPO; ECTP;	
XX	XX	XX	KW anaemia; therapy; antianaemic.	
AC	AD079063;	OS Homo sapiens.	OS Synthetic.	
XX	XX	XX	OS Homo sapiens.	
DT	29-JUL-2004 (first entry)	XX	Key location/Qualifiers	
XX	XX	XX	FT 1. .192	
DE	Human thrombopoietin/erythropoietin fusion protein #2.	XX	FT /note= "human erythropoietin"	
XX	XX	XX	FT 193. .220	
KW	fusion protein; carboxy terminal peptide; CTP; human; thrombopoietin;	XX	FT /note= "HCG beta subunit CTP"	
TPO; erythropoietin; EPO; anaemia.	XX	XX	PN WO200249194-A1.	
OS	Chimeric.	XX	PN 20-JUN-2002.	
XX	XX	XX	PP 10-DEC-2001; 2001WO-KR002137.	
PN	GB2002580-A.	XX	PR 11-DEC-2000; 2000KR-00075230.	
XX	XX	XX	PR 21-NOV-2001; 2001KR-00072713.	
PD	04-JUN-2003.	XX	PA (CHRI-) CHEIL JEDANG CO.	
XX	XX	XX	XX	
PR	06-AUG-2002; 2002GB-00018252.	XX	PI Lee D., Oh M., Kim K., Chung B., Ha B., Park J.;	
XX	XX	XX	DR N-FSDB; ABQ81360.	
PA	(CHRI-) CHEIL JEDANG CORP.	XX	PT WPI; 2002-713247/77.	
XX	XX	XX	DR	
PA	Lee D., Oh M., Chung B., Park J., Kim K.	XX	N-FSDB; ABQ81360.	
XX	XX	XX	PT Novel fusion protein useful for industrial purposes, comprises carboxy terminal of human erythropoietin fused with carboxy terminal of beta subunit of human chorionic gonadotropin.	
PS	Novel fusion protein having enhanced in vivo activity useful for treating anaemia, comprises carboxy terminal peptide of thrombopoietin fused with carboxy terminal of human erythropoietin.	XX	PT fragment of beta subunit of human chorionic gonadotropin.	
XX	XX	XX	PS Example 1; Fig 2; 30PP; English.	
PT	The invention comprises a fusion protein consisting of the carboxy terminal peptide (CTP) of human thrombopoietin (TPO) fused to the carboxy terminal of human erythropoietin (EPO). The fusion protein of the invention is useful for the treatment of anaemia. The present amino acid sequence represents a human thrombopoietin/erythropoietin fusion protein of the invention.	XX	CC The present sequence is the protein sequence of a fusion protein, termed ECTP, in which the C-terminus of human erythropoietin (EPO) is fused with a C-terminal peptide (CTP) (see also ABB81359) of of human chorionic gonadotrophin (HCG) beta subunit. The CTP comprises amino acids 118-145 (see also ABB79937) of the HCG beta subunit. The invention provides ECTP fusion protein and nucleotide sequences encoding it, a plasmid containing the nucleotide sequences, a host cell (e.g. CHO) transfected with the plasmid, and a method for producing the fusion protein by cultivation of the transfected cell line. Fusion to HCG beta subunit CTP enhances the in vivo activity of EPO for treatment of anaemia. The CTP provides extra glycosylation sites, increasing the half-life of EPO without causing any loss of the inherent activity of EPO without causing any antigenicity when applied to the human body. Pharmacokinetic experiments performed in mice showed that ECTP had 2.5 times longer half-life than EPO	
SQ	Sequence 209 AA;	XX	CC Sequence 220 AA;	
QY	Query Match 100.0%; Score 846; DB 7; Length 209; Best Local Similarity 100.0%; Pred. No. 3.1e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	XX	Query Match 100.0%; Score 846; DB 5; Length 220; Best Local Similarity 100.0%; Pred. No. 3.4e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	
Db	1 APPRLICDSRVLERYLEAKEAENITTCGAEHCSENNTIVPDTCVNFTYAWKRMVGQA 60	XX	Db 1 APPRLICDSRVLERYLEAKEAENITTCGAEHCSENNTIVPDTCVNFTYAWKRMVGQA 60	
QY	2 APPRLICDSRVLERYLEAKEAENITTCGAEHCSENNTIVPDTCVNFTYAWKRMVGQA 87	XX	QY 1 APPRLICDSRVLERYLEAKEAENITTCGAEHCSENNTIVPDTCVNFTYAWKRMVGQA 60	
Db	61 VEVNQGLALISEAVRQGQALLVNSQPMWPLQHDKAVSGRLSITLRAQKEAIS 120	XX	Db 28 APPRLICDSRVLERYLEAKEAENITTCGAEHCSENNTIVPDTCVNFTYAWKRMVGQA 87	
QY	88 VEVNQGLALISEAVRQGQALLVNSQPMWPLQHDKAVSGRLSITLRAQKEAIS 147	XX	QY 61 VEVNQGLALISEAVRQGQALLVNSQPMWPLQHDKAVSGRLSITLRAQKEAIS 120	
Db	121 PPDAASAAPTPITTDFTFKLFRVSNFLRGKLYKTYGACRTGD 165	XX	Db 148 PPDAASAAPLRTTADTFRKLFRVSNFLRGKLYKTYGACRTGD 192	
QY	148 PPDAASAAPLRTTADTFRKLFRVSNFLRGKLYKTYGACRTGD 192	XX	QY 121 PPDAASAAPLRTTADTFRKLFRVSNFLRGKLYKTYGACRTGD 165	

			DE	Recombinant hematopoietic molecule 1.
RESULT 101			XX	IL-3; EPO; haematopoiesis.
ABR57656			KW	
ID	ABR57656	standard; protein; 220 AA.	OS	Homo sapiens.
XX			XX	
AC	ABR57656;		AC	W09206116-A.
XX			PD	16-APR-1992.
DT	04-DEC-2003	(first entry)	XX	
DE	Fusion protein comprising erythropoietin and mutant CTP fragment.		PF	26-SEP-1991; 91WO-US007053.
XX			XX	
KW	Antianemic; human; EPO; CTP; HCG; erythropoietin;		PR	28-SEP-1990; 90US-0058958.
Carboxyl Terminal Peptide; human chorionic gonadotropin; anaemia.			PA	(ORTH) ORTHO PHARM CORP.
XX			XX	
OS	Synthetic.		PT	Rosen, JT;
XX			XX	
PN	EP1316561-A1.		DR	WPI; 1992-150819/18.
XX			XX	
PD	04-JUN-2003.		PT	Recombinant haematopoietic molecules useful in treating anaemia(s) -
XX			PT	comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
PP	14-AUG-2002; 2002EP-00255679.		PT	later myeloid differentiation activity.
XX			XX	
PR	03-DEC-2001; 2001KR-00075994.		PS	Disclosure; Page 34; 82pp; English.
XX			XX	
PA	(CHEI-) CHEIL JEDANG CORP.		CC	This protein sequence given comprises the entire amino acid sequence of a
XX			CC	recombinant haematopoietic molecule, with the amino portion comprising IL-
PI	Lee D, Oh M, Kim K, Chung B, Park J;		CC	3 and the carboxy portion comprising EPO. (Specific sequences for these
XX			CC	portions are given in AAR3591 and AAR23593.) Within the scope of the
DR	WPI; 2003-495240/47.		CC	invention hybrid molecules were produced which contain at least a portion
DR	N-P5DB; ACC80208.		CC	of an early MDP and at least a portion of a late MDP covalently linked.
XX			CC	These compounds can be used to promote hematopoiesis in patient. The
PT	New fusion protein, useful for treating anaemia, comprises human erythropoietin having a carboxyl terminal and a carboxyl terminal peptide		CC	bonding of the early and late factors allows a very high conc. of late
PT	fragment of a human chorionic gonadotropin beta-subunit linked to the		CC	MDP at the surface of a cell which the early MDP is bound. It also allows
PT	carboxyl terminal.		CC	the early MDP to act more specifically to stimulate only the desired
XX			CC	lineage, thus reducing undesirable effects. These compounds are useful
PS	Disclosure; Page 8-9; 19pp; English.		CC	for treating anaemias of various origins eg. renal failure and AIDS. It is
XX			CC	easier to produce and administer one recombinant molecule rather than two
CC	human erythropoietin (EPO) and a mutant of a Carboxyl Terminal Peptide (CTP; ABR57655) fragment of a human chorionic gonadotropin (HCG) beta-		CC	separate molecules
CC	subunit with 1-4 amino acid substitutions in the CTP fragment. The fusion		XX	
CC	protein is useful in preparing a medicament for treating anaemia		SQ	Sequence 302 AA;
XX				
SQ	Sequence 220 AA;			
Query	Match	100.0%; Score 846; DB 7; Length 220;	Query	Match
Best Local Similarity	100.0%; Pred. No. 3, 4e-86;	100.0%; Score 846; DB 2; Length 302;	Best Local Similarity	100.0%; Pred. No. 5, 3e-86;
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	Mismatches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	
QY	1 APPRLICDSRVLEVLLEAKEAENITTCGAECNSLNENITTVDPDKVNYAWGRMEVQQA	QY	1 APPRLICDSRVLEVLLEAKEAENITTCGAECNSLNENITTVDPDKVNYAWGRMEVQQA	
Db	60	Db	60	
QY	2 APPRLICDSRVLEVLLEAKEAENITTCGAECNSLNENITTVDPDKVNYAWGRMEVQQA	QY	61 VEWQGLALLSEAVLRGQALLNSQSOPWEPLQLAVDVKAVSGIRSLSLTILRAAGQKEAIS	
Db	87	Db	120	
QY	61 VEWQGLALLSEAVLRGQALLNSQSOPWEPLQLAVDVKAVSGIRSLSLTILRAAGQKEAIS	QY	197 VEWQGLALLSEAVLRGQALLNSQSOPWEPLQLAVDVKAVSGIRSLSLTILRAAGQKEAIS	
Db	147	Db	256	
QY	121 PPDAAASAPLRTTADTRKLRFLRVSNFLRGKLUYGEACTGD 165	QY	121 PPDAAASAPLRTTADTRKLRFLRVSNFLRGKLUYGEACTGD 165	
Db	148 PPDAAASAPLRTTADTRKLRFLRVSNFLRGKLUYGEACTGD 192	Db	257 PPDAAASAPLRTTADTRKLRFLRVSNFLRGKLUYGEACTGD 301	
RESULT 102				
AAR23596			RESULT 103	
ID	AAR23598 standard; protein; 303 AA.		AAR23598	
XX			ID	AAR23598 standard; protein; 303 AA.
AC	AAR23598;		XX	
XX			AC	AAR23598;
DT	20-OCT-1992 (first entry)		XX	
XX			DT	20-OCT-1992 (first entry)
DE	Recombinant hematopoietic molecule 3.		XX	Recombinant hematopoietic molecule 3.
XX			XX	IL-3; EPO; haematopoiesis.
XX			OS	Homo sapiens.
XX				

XX
PN WO9206116-A.
XX PN
PD 16-APR-1992.
XX
XX 26-SEP-1991; 91WO-US007053.
PF XX
PR 28-SEP-1990; 90US-00589958.
PA (ORTH) ORTHO PHARM CORP.
XX
PI Rosen JI;
XX DR
DR WPI; 1992-150819/18.
XX PT
PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
PT later myeloid differentiation activity.
XX PS
PS Disclosure; Page 38; 82pp; English.

XX CC
CC This protein sequence given comprises the entire amino acid sequence of a recombinant haematopoietic molecule, with the amino portion comprising EPO and the carboxyl portion comprising IL-3. (Specific sequences for these portions are given in AAR23591 and AAR23593.) Within the scope of the invention hybrid molecules were produced which contain at least a portion of an early MDF and at least a portion of a late MDF covalently linked. These compounds can be used to promote hematopoiesis in a patient. The bonding of the early and late factors allows a very high conc. of late MDF at the surface of a cell which the early MDF is bound. It also allows the early MDF to act more specifically to stimulate only the desired lineage, thus reducing undesirable effects. These compounds are useful for treating anaemias of various origins eg. renal failure and AIDS. It is easier to produce and administer one recombinant molecule rather than two separate molecules

XX SQ
Sequence 303 AA;

Query Match	Score	DB	Length
Best Local Similarity	100.0%	2;	303;
Matches	165;	Pred. No.	5.4e-86;
Conservative	0;	Mismatches	0;
Indels	0;	Gaps	0;

CC 1 APPRLICDSVRVLERYLLEAKEAEINTTGCACRHSCLNENITVPDKVNYAWKRMEVGQOA 60
Db 1 APPRLICDSVRVLERYLLEAKEAEINTTGCACRHSCLNENITVPDKVNYAWKRMEVGQOA 60
QY 61 VEWVOGLALISEAVURGOALLVNSQWPMLQPLQHVDKAVSGLRSLTLLRAGQKEAIS 120
Db 61 VEWVOGLALISEAVURGOALLVNSQWPMLQPLQHVDKAVSGLRSLTLLRAGQKEAIS 120
QY 121 PPDAAASAAPRTTADTPRLFLRVISNFRLGRKLYTGACRTGD 165
Db 121 PPDAAASAAPRTTADTPRLFLRVISNFRLGRKLYTGACRTGD 165
Db 276 PPDAAASAAPRTTADTPRLFLRVISNFRLGRKLYTGACRTGD 320

RESULT 104
AAR23075
ID AAR23075 standard; protein; 321 AA.
XX
AC AAR23075:
XX DT 20-OCT-1992 (first entry)
XX DE IL-3:Epo short, recombinant hematopoietic molecule.
XX KW Early MDF; late MDF; haematopoiesis; IL-3; Epo; growth factor.
XX OS Homo sapiens.
XX FH Key
FT Peptide 1:-19
FT /label= sig_peptide 20 . .321
FT /label= mat_protein

XX
PN WO9206116-A.
XX PN
PD 16-APR-1992.
XX
XX 26-SEP-1991; 91WO-US007053.
PF XX
PR 28-SEP-1990; 90US-00589958.
PA (ORTH) ORTHO PHARM CORP.
XX
PI Rosen JI;
XX DR
DR WPI; 1992-150819/18.
XX PT
PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
PT later myeloid differentiation activity.
XX PS
PS Disclosure; Page 42; 82pp; English.

XX CC
CC The amino acid sequence given is an IL-3:Epo hybrid growth factor derived from a construction formed by ligating various synthetic oligonucleotides corresponding to Epo and IL-3 gene sequences. This hybrid growth factor is a recombinant haematopoietic molecule which contains at least a portion of an early MDF and at least a portion of a late MDF covalently linked. This compound can be used to promote hematopoiesis in a patient. The bonding of the early and late factors allows a very high conc. of late MDF at the surface of a cell which the early MDF is bound. It also allows the early MDF to act more specifically to stimulate only the desired lineage, thus reducing undesirable effects. These compounds are useful for treating anaemias of various origins eg. renal failure and AIDS. It is easier to produce and administer one recombinant molecule rather than two separate molecules

XX SQ
Sequence 321 AA;

Query Match	Score	DB	Length
Best Local Similarity	100.0%	2;	321;
Matches	165;	Pred. No.	5.8e-86;
Conservative	0;	Mismatches	0;
Indels	0;	Gaps	0;

QY 1 APPRLICDSVRVLERYLLEAKEAEINTTGCACRHSCLNENITVPDKVNYAWKRMEVGQOA 60
Db 156 APPRLICDSVRVLERYLLEAKEAEINTTGCACRHSCLNENITVPDKVNYAWKRMEVGQOA 215
QY 61 VEWVOGLALISEAVURGOALLVNSQWPMLQPLQHVDKAVSGLRSLTLLRAGQKEAIS 120
Db 216 VEWVOGLALISEAVURGOALLVNSQWPMLQPLQHVDKAVSGLRSLTLLRAGQKEAIS 275
QY 121 PPDAAASAAPRTTADTPRLFLRVISNFRLGRKLYTGACRTGD 165
Db 276 PPDAAASAAPRTTADTPRLFLRVISNFRLGRKLYTGACRTGD 320

RESULT 105
AAR23597
ID AAR23597 standard; protein; 321 AA.
XX AC AAR23597:
XX DT 20-OCT-1992 (first entry)
XX DE Recombinant hematopoietic molecule 2.
XX KW IL-3; EPO; haematopoiesis.
XX OS Homo sapiens.
XX FN WO9206116-A.
XX PD 16-APR-1992.

XX DR Rosen JI;
 PR XX WPI; 1992-150819/18.
 XX DR
 XX PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
 PA PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
 PI PT later myeloid differentiation activity.
 XX PS Disclosure; Page 36; 82pp; English.
 XX PS Disclosure; Page 39; 82pp; English.
 XX CC Recombinant haematopoietic molecule, with the amino portion comprising IL-
 CC recombinant haematopoietic molecule, with the amino portion comprising IL-
 CC 3 and the carboxy portion comprising EPO. (Specific sequences for these
 CC portions are given in AAR23591 and AAR23593.) Within the scope of the
 CC invention hybrid molecules were produced which contain at least a portion
 CC of an early MDP and at least a portion of a late MDP covalently linked.
 CC These compounds can be used to promote hematopoiesis in a patient. The
 CC bonding of the early and late factors allows a very high conc. of late
 CC MDP at the surface of a cell which the early MDP is bound. It also allows
 CC the early MDP to act more specifically to stimulate only the desired
 CC lineage, thus reducing undesirable effects. These compounds are useful
 CC for treating anaemias of various origins eg. renal failure and AIDS. It is
 CC easier to produce and administer one recombinant molecule rather than two
 CC separate molecules
 XX SQ Sequence 321 AA;
 Query Match 100.0%; Score 846; DB 2; Length 321;
 Best Local Similarity 100.0%; Pred. No. 5.8e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 APPRLICDSRVLERYLEAKRENTITGCAHCSLENITVPDTKVNPFYAKWKRMEVQQA 60
 Db 156 APPRLICDSRVLERYLEAKRENTITGCAHCSLENITVPDTKVNPFYAKWKRMEVQQA 215
 QY 61 VEWQGLALLSEAVRQQLVNSQPWEPLQHDKAVSGLSLTTLRAIGAQEAIIS 120
 Db 216 VEWQGLALLSEAVRQQLVNSQPWEPLQHDKAVSGLSLTTLRAIGAQEAIIS 275
 QY 121 PPDAASAPLRTITADTFRKLFRVVSNFLRGKLKLYGEACTGD 165
 Db 276 PPDAASAPLRTITADTFRKLFRVVSNFLRGKLKLYGEACTGD 320
 RESULT 106
 AAR23599 ID AAR23599 standard; protein; 322 AA.
 AC AAR23599;
 XX DT 20-OCT-1992 (first entry)
 DE Recombinant hematopoietic molecule 4.
 XX KW IL-3; EPO; haematopoiesis.
 OS Homo sapiens.
 PN WO2006116-A.
 PD 16-APR-1992.
 PR 26-SEP-1991; 91WO-US007053.
 PR 28-SEP-1990; 90US-00589958.
 PA (ORTH) ORTHO PHARM CORP.
 XX DR Rosen JI;
 PR XX WPI; 1992-150819/18.
 XX DR
 XX PT Recombinant haematopoietic molecules useful in treating anaemia(s) -
 PA PT comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
 PI PT later myeloid differentiation activity.
 XX PS Disclosure; Page 36; 82pp; English.
 XX PS Disclosure; Page 39; 82pp; English.
 XX CC Recombinant haematopoietic molecule, with the amino portion comprising EPO
 CC and the carboxy portion comprising IL-3. (Specific sequences for these
 CC portions are given in AAR23591 and AAR23593.) Within the scope of the
 CC invention hybrid molecules were produced which contain at least a portion
 CC of an early MDP and at least a portion of a late MDP covalently linked.
 CC These compounds can be used to promote hematopoiesis in a patient. The
 CC bonding of the early and late factors allows a very high conc. of late
 CC MDP at the surface of a cell which the early MDP is bound. It also allows
 CC the early MDP to act more specifically to stimulate only the desired
 CC lineage, thus reducing undesirable effects. These compounds are useful
 CC for treating anaemias of various origins eg. renal failure and AIDS. It is
 CC easier to produce and administer one recombinant molecule rather than two
 CC separate molecules
 XX SQ Sequence 322 AA;
 Query Match 100.0%; Score 846; DB 2; Length 322;
 Best Local Similarity 100.0%; Pred. No. 5.9e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 QY 1 APPRLICDSRVLERYLEAKRENTITGCAHCSLENITVPDTKVNPFYAKWKRMEVQQA 60
 Db 1 APPRLICDSRVLERYLEAKRENTITGCAHCSLENITVPDTKVNPFYAKWKRMEVQQA 60
 QY 61 VEWQGLALLSEAVRQQLVNSQPWEPLQHDKAVSGLSLTTLRAIGAQEAIIS 120
 Db 61 VEWQGLALLSEAVRQQLVNSQPWEPLQHDKAVSGLSLTTLRAIGAQEAIIS 120
 QY 121 PPDAASAPLRTITADTFRKLFRVVSNFLRGKLKLYGEACTGD 165
 Db 121 PPDAASAPLRTITADTFRKLFRVVSNFLRGKLKLYGEACTGD 165

PA	XX	(ORTH) ORTHO PHARM CORP.	
PT	XX	Rosen JI;	
XX	XX		
DR	XX		
N-PSDB;	AAQ24282.		
PT	XX	Recombinant haematopoietic molecules useful in treating anaemia(s) -	
PT	XX	comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and	
PT	XX	later myeloid differentiation activity.	
PS	XX		
Disclosure; Page 44; 82pp; English.			
XX	CC	The amino acid sequence given is an Epo:IL-3 hybrid growth factor derived	
CC	CC	from a construction formed by ligating the native Epo signal sequence and	
CC	CC	various synthetic oligonucleotides corresponding to Epo and IL-3 gene	
CC	CC	sequences. This hybrid growth factor is a haematopoietic molecule which	
CC	CC	contains at least a portion of an early MDF and at least a portion of a	
CC	CC	late MDF covalently linked. This compound can be used to promote	
CC	CC	haematopoiesis in a patient. The bonding of the early and late factors	
CC	CC	allows a very high conc. of late MDF at the surface of a cell which the	
CC	CC	early MDF is bound. It also allows the early MDF to act more specifically	
CC	CC	to stimulate only the desired lineage, thus reducing undesirable effects.	
CC	CC	These compounds are useful for treating anaemias of various origins	
CC	CC	eg. renal failure and AIDS. It is easier to produce and administer one	
CC	CC	recombinant molecule rather than two separate molecules	
XX	CC	recombinant molecule rather than two separate	
SQ	XX	Sequence 330 AA;	
Query Match	100.0%; Score 846; DB 2; Length 330;		
Best Local Similarity	100.0%; Pred. No. 6.1e-86;		
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		
Oy	1 APPRLICDSVRLEYLEAKEAENITTCGAHCISLENINITYPDTKUNFYAWKRMEVGQQA 60		
Db	28 APPRLICDSVRLEYLEAKEAENITTCGAHCISLENINITYPDTKUNFYAWKRMEVGQQA 87		
Oy	61 VEWQGALLSEAVRQALINNSQWEPLQHVDKAVSGRLSLTTLRAAGQKAIS 120		
Db	88 VEWQGALLSEAVRQALINNSQWEPLQHVDKAVSGRLSLTTLRAAGQKAIS 147		
Oy	121 PPDRASAAPLRTITADTPFRKLFRVYSNTRGKLYTGEACRTGD 165		
Db	148 PPDRASAAPLRTITADTPFRKLFRVYSNTRGKLYTGEACRTGD 192		
RESULT 108			
AAR23078			
ID	AAR23078 standard; protein; 340 AA..		
XX			
AC	AAR23078;		
XX			
DT	20-OCT-1992 (first entry)		
XX			
DE	IL-3:Epo Flex, recombinant hematopoietic molecule.		
XX			
KW	Early MDF; late MDF; haematopoiesis; IL-3; Epo; growth factor; linker.		
OS	Homo sapiens.		
XX			
FR	Location/Qualifiers		
PT	Key Peptide 1..19 /label= sig_peptide		
FT	Protein 20..339 /label= mat_protein		
XX			
PN	WO9206116-A.		
XX			
PD	16-APR-1992.		
XX			
PP	26-SEP-1991; 91WO-US007053.		
XX			
PR	28-SEP-1990; 90US-00589958.		
PS	XX	Recombinant haematopoietic molecules useful in treating anaemia(s) -	
PS	XX	comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and	
PS	XX	later myeloid differentiation activity.	
PS	XX	Disclosure; Page 49; 82pp; English.	
SQ	XX	Sequence 340 AA;	
Query Match	100.0%; Score 846; DB 2; Length 340;		
Best Local Similarity	100.0%; Pred. No. 6.3e-86;		
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;		
Oy	1 APPRLICDSVRLEYLEAKEAENITTCGAHCISLENINITYPDTKUNFYAWKRMEVGQQA 60		
Db	175 APPRLICDSVRLEYLEAKEAENITTCGAHCISLENINITYPDTKUNFYAWKRMEVGQQA 234		
Oy	61 VEWQGALLSEAVRQALINNSQWEPLQHVDKAVSGRLSLTTLRAAGQKAIS 120		
Db	235 VEWQGALLSEAVRQALINNSQWEPLQHVDKAVSGRLSLTTLRAAGQKAIS 294		
Oy	121 PPDRASAAPLRTITADTPFRKLFRVYSNTRGKLYTGEACRTGD 165		
Db	295 PPDRASAAPLRTITADTPFRKLFRVYSNTRGKLYTGEACRTGD 339		
RESULT 109			
AAR23079			
ID	AAR23079 standard; protein; 349 AA..		
XX			
AC	AAR23079;		
XX			
DT	20-OCT-1992 (first entry)		
XX			
DB	Epo:IL-3 Flex, recombinant hematopoietic molecule.		
XX			
KW	Early MDF; late MDF; haematopoiesis; Epo; IL-3; linker; growth factor.		
OS	Homo sapiens.		
XX			
FT	Location/Qualifiers		
FT	Key Peptide 1..27 /label= sig_peptide		
FT	Protein 28..349 /label= mat_protein		
XX			
PN	WO9206116-A.		
XX			
PD	16-APR-1992.		
XX			

PF	29-NOV-2001; 2001KR-00074975.
XX	
PR	(CHERI-) CHEIL JEDANG CORP.
XX	
PA	(ORTH) ORTHO PHARM CORP.
XX	
PI	Rosen JI;
XX	
DR	WPI; 1992-150819/18.
XX	
DR	N-PSDB; AAQ24285.
XX	
PT	Recombinant haematopoietic molecules useful in treating anaemia (s) -
PT	comprise IL3 or GM-CSF and EPO, G-CSF, IL-5 or M-CSF and has early and
PT	later myeloid differentiation activity.
XX	
PS	Disclosure; Page 51; 82pp; English.
XX	
CC	The amino acid sequence given is an Epo:IL-3 hybrid growth factor derived
CC	from a construction formed by ligating the native Epo signal sequence and
CC	various synthetic oligonucleotides corresponding to Epo and IL-3 gene
CC	sequences. This molecule is comparable to the sequence given in ABR23076
CC	and contains a flexible linker molecule. This hybrid growth factor is a
CC	haemopoietic molecule which contains at least a portion of an early MDP
CC	and at least a portion of a late MDP covalently linked. This compound can
CC	be used to promote hematopoiesis in a patient. The bonding of the early
CC	and late factors allows a very high conc. of late MDP at the surface of a
CC	cell which the early MDP is bound. It also allows the early MDP to act
CC	more specifically to stimulate only the desired lineage, thus reducing
CC	undesirable effects. These compounds are useful for treating anaemias of
CC	various origins e.g. renal failure and AIDS. It is easier to produce and
CC	administer one recombinant molecule rather than two separate molecules
XX	
SQ	Sequence 349 AA;
	Best Local Similarity 100.0%; Score 846; DB 2; Length 349;
Matches	165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Oy	1 APPRLICDSRVLYRVLLEAKEAENITGCAEHCSLNENITPDTKNFYAWKRMEMQQA 60
Db	28 APPRLICDSRVLYRVLLEAKEAENITGCAEHCSLNENITPDTKNFYAWKRMEMQQA 87
Oy	61 VEWMOGLALLSEAVLVRGQALLNSSQPEPLQLHVDRKAVSGRSLTTLRAKGKAIS 120
Db	88 VEWMOGLALLSEAVLVRGQALLNSSQPEPLQLHVDRKAVSGRSLTTLRAKGKAIS 147
Oy	121 PPDAAASAPLRTTADTPRKLFVYSNFLRGKUKLYGEACRTGD 165
Db	148 PPDAAASAPLRTTADTPRKLFVYSNFLRGKUKLYGEACRTGD 192
	RESULT 111
	AAW99360
ID	AAW99360 standard; protein; 376 AA.
XX	
AC	AAW99360;
XX	
DT	21-MAY-1999 (first entry)
XX	
DE	Human erythropoietin homodimer fusion protein.
KW	Human; erythropoietin; dimer; trimer; polymer; fusion protein; cancer;
KW	biological activity; anaemia; proliferation; differentiation; progenitor;
KW	leucocyte; granulocyte; blood; myelosuppressed patient.
XX	
OS	Homo sapiens.
OS	Synthetic.
PN	W09902710-A1.
XX	
DD	21-JAN-1999.
XX	
PF	09-JUL-1998; 98WO-US013944.
XX	
PR	10-JUL-1997; 97US-0089929.
PR	03-FEB-1998; 98US-00018138.
XX	
PA	(BETH-) BETH ISRAEL DEACONESS MEDICAL CENT.
XX	
PI	Sytkowski AJ;
XX	
DR	WPI; 1999-120911/10.
XX	
DR	N-PSDB; AAQ25701.

RESULT 110

ADO7962 standard; protein; 370 AA.

ID ADO7962

standard; protein; 370 AA.

ADO7962;

29-JUL-2004 (first entry)

Human thrombopoietin/erythropoietin fusion protein #1.

fusion protein; carboxy terminal peptide; CTP; human; thrombopoietin;

TPO; erythropoietin; EPO; anaemia.

Homo sapiens.

Chimeric.

GB2382580-A.

04-JUN-2003.

XX

06-AUG-2002; 2002GB-00018252.

XX
PT New fusion protein with increased activity comprising at least two
protein molecules - used to, e.g. treat erythropoietin related deficiency
PT states for treatment of anaemia.
XX
PS Example 1; FIG 16a-C; 119pp; English.
XX
CC This sequence represents a human erythropoietin (EPO) homodimeric fusion
protein. The invention relates to the production of dimeric, trimeric or
polymeric fusion proteins with increased biological activity. The fusion
proteins are used to treat or prevent protein-related deficiency states,
specifically, where the protein is erythropoietin (EPO; AAX2589),
CC anaemia, but also for increasing proliferation, differentiation and
activity of haematopoietic progenitors (e.g. increasing numbers of
leucocytes and granulocytes in the blood of myelosuppressed patients) or
CC for treating cancer and other cell growth disorders
XX
SQ Sequence 376 AA;

Query Match Score 846; DB 2; Length 376;
Best Local Similarity 100.0%; Pred. No. 7, 9e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTCGAECHECSINLENITIVPDTKVNFTAWKMERVGQA 60
Db 28 APPRLICDSRVLERYLLEAKEAENITTCGAECHECSINLENITIVPDTKVNFTAWKMERVGQA 87

QY 61 VEWVOGLALIISEAVERGQQLVNSQPMPLQLQHDKAVSGLSLTLLRAKGKEAIS 120
Db 88 VEWVOGLALIISEAVERGQQLVNSQPMPLQLQHDKAVSGLSLTLLRAKGKEAIS 147

QY 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRGD 165
Db 148 PPDASAAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRGD 192

RESULT 112
AEB12283 ID AEB12283 standard; protein; 397 AA.
XX AC AEB12283;
XX DT 22-SEP-2005 (first entry)
DB Human IgG2- erythropoietin fusion protein huFcg2h (FN-AQ) -M1-EPO.

XX KW Erythropoietin; muttein; fusion protein; protein therapy; antianemic;
hematological disease; immunoglobulin.
XX OS Homo sapiens.
OS Synthetic.
PN WO2005063808-A1.
XX PD 14-JUL-2005.
XX PF 22-DEC-2004; 2004WO-EP014608.
PR 31-DEC-2003; 2003US-0533858P.
XX PA (MERCK) MERCK PATENT GMBH.
XX PI Gillies SD, Lauder S;
DR WPI; 2005-506648/51.

PT New purified dimeric fusion protein comprises a dimeric Fc portion of a
human immunoglobulin G molecule and human erythropoietin, for treating
hematopoietic disorders or deficiencies in a mammal.

PS Claim 14; SEQ ID NO 14; 87pp; English.
CC The invention relates to a purified dimeric fusion protein comprising a

CC dimeric Fc portion of a human immunoglobulin (Ig)G molecule comprising a
hinge region, a CH2, and a CH3 domain, and human erythropoietin (EPO),
CC where each chain of the dimeric Fc portion is linked via its C-terminus
CC directly or via a linker peptide to the N-terminus of an EPO molecule.
The molecule is highly sialylated by comprising 15-28 sialic acid
CC residues, the CH2 domain derives from human IgG2 (and is modified by
CC replacing the amino acid residue Phe and Asn within the Gln-Phe-Asn-Ser
CC sequence track of the CH2 domain with Ala and Asn, thus forming the
CC amino acid sequence track near the C-terminus of the CH3 domain is
CC replaced with Ala-Thr-Ala-Thr). Also included are a DNA molecule encoding
the fusion protein, a pharmaceutical composition for the treatment of
hematopoietic disorders or deficiencies in a mammal (comprising an amount
CC of the Fc-EPO fusion protein, optionally together with a pharmaceutical
CC carrier, diluent, or excipient), a population of purified highly
sialylated Fc-PO fusion proteins for administration to a mammal (the Fc-
CC EPO fusion proteins comprising an Fc portion towards the N-terminus of
the Fc-EPO fusion proteins, a method of producing a
CC population of highly sialylated purified recombinant Fc-EPO fusion
proteins and a method of selecting a baby hamster kidney (BHK) cell
CC stably maintaining a nucleic acid sequence encoding an Fc-EPO fusion
protein. The purified dimeric fusion protein is useful for treating
hematopoietic disorders (hematological disease) or deficiencies in a
mammal. The present sequence represents the Human IgG2- erythropoietin
CC Fusion protein of the invention, huFcg2h (FN-AQ)-M1-EPO (carrying an FN to
AQ mutation in the IgG2 CH2 domain, eliminating a T-cell epitope/N-
CC glycosylation site).

SQ Sequence 397 AA;

Query Match Score 846; DB 9; Length 397;
Best Local Similarity 100.0%; Pred. No. 7, 9e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTCGAECHECSINLENITIVPDTKVNFTAWKMERVGQA 60
Db 232 APPRLICDSRVLERYLLEAKEAENITTCGAECHECSINLENITIVPDTKVNFTAWKMERVGQA 291

QY 61 VEWVOGLALIISEAVERGQQLVNSQPMPLQLQHDKAVSGLSLTLLRAKGKEAIS 120
Db 292 VEWVOGLALIISEAVERGQQLVNSQPMPLQLQHDKAVSGLSLTLLRAKGKEAIS 351

QY 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRGD 165
Db 352 PPDASAAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRGD 396

RESULT 113
ABU64200 ID ABU64200 standard; protein; 428 AA.

XX AC ABU64200;
XX DT 11-MAR-2004 (first entry)
XX DE Plasmid PBD-dC-natEpoFc nativeEpo/Fcgamma1 insert protein.

XX KW Transepithelial systemic delivery; therapeutic delivery; aerosol;
KW FcRn binding partner; lung.

XX OS Synthetic.
XX PN WO2003077834-A2.

XX PD 25-SEP-2003.
XX PR 03-JUL-2002; 2002WO-US021335.

XX PR 15-MAR-2002; 2002US-0364482P.

XX PA (BGH) BRIGHAM & WOMENS HOSPITAL INC.

PT	Blumberg RS, Lencer WI, Simister NE, Bitonti AJ;	FT /note= "Human mature EPO"	
XX		FT 194. .201	
DR	WPI; 2003-767442/72.	FT /note= "8 residue peptide linker (SEQ ID NO:2)"	
DR	N-PSDB; AU56123.	FT 202. .428	
XX		FT /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
PT	Aerosol useful for systemic delivery of a therapeutic agent e.g., erythropoietin, growth hormone, interferon-alpha, or interferon-beta, comprises a conjugate of the agent and neonatal epithelial receptor-binding partner.	IX	
PT		PN WO2004004798-A2.	
PT		PD 15-JAN-2004.	
XX		XX	
PS	Example 5; FIG 5B; DPP; English.	PF 09-MAY-2003; 2003WO-US014428.	
XX		XX	
CC	The present invention relates to an aerosol which comprises a conjugate of a therapeutic agent and neonatal Fc receptor (FcRn) binding partner.	PR 03-JUL-2002; 2002WO-US021335.	
CC	The particles in the aerosol have a mass median aerodynamic diameter (MMAD) of at least 3 micro m. The aerosol can be used for the systemic delivery of a therapeutic agent (e.g. antigen (e.g. tumour antigen), polypeptide, oligonucleotide (e.g. antisense oligonucleotide), erythropoietin, growth hormone, interferon-alpha, interferon-beta and follicle stimulating hormone). The present sequence is a protein used in the exemplification of the invention	XX	
CC		(BGHM) BRIGHAM & WOMENS HOSPITAL INC.	
CC		PA (UYBR-) UNIV BRANDEIS.	
CC		PA (CHIL-) CHILDRENS MEDICAL CENT.	
CC		PA (SYNT-) SYNTONIX PHARM INC.	
CC		XX	
CC		Blumberg RS, Lencer WI, Simister NE, Bitonti AJ;	
CC		XX	
XX		DR WPI; 2004-099348/10.	
SQ	Sequence 428 AA;	DR N-PSDB; ADO10512.	
Query	Match 100.0%; Score 846; DB 7; Length 428;	XX	
Best Local Similarity 100.0%; Pred. No. 8. 9e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	PR Systemic delivery of therapeutic agent involves administering effective amount of aerosol of therapeutic agent and neonatal Fc receptor (FcRn) binding partner to lung.		
Qy 1 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 60	PT Region	Region /note= "Human mature EPO"	
Db 28 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 87	FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"	
Qy 61 VEVWQGLALLSEAVLQGQLNSSOWEPOLHVDAVSGRSLSLTIRALGAQKRAIS 120	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Db 88 VEVWQGLALLSEAVLQGQLNSSOWEPOLHVDAVSGRSLSLTIRALGAQKRAIS 147	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Qy 121 PPDAAASAPURITATDPRKURVYSNPLRKSLKLTYGCAARTGTD 165	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Db 148 PPDAAASAPURITATDPRKURVYSNPLRKSLKLTYGCAARTGTD 192	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
RESULT 114		FT Region	Region /note= "Human mature EPO"
ADO10513		FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"
ID ADO10513 standard; protein; 428 AA.		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
XX		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
AC ADO10513;		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
XX		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
DT 01-JUL-2004 (first entry)		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
XX		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
DE EPO signal peptide/EPO/IgG1 Fc fragment fusion protein, SEQ ID NO:10.		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
XX		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
Drug delivery; aerosol; transepithelial; FcRn ligand; neonatal Fc receptor; central airway epithelium; lung; antigen; tumour antigen; erythropoietin; EPO; growth hormone; interferon-alpha; IFN-alpha; interferon-beta; FSH; therapeutic antibody; CAMPATH; SIMULECT; ZENPAX; SYNAGIS; RITUXAN; HERCEPTIN; REMICADE; HUMIRA; CEA-CIDE. Therapeutics administered using the method of the invention may be used to treat deep lung diseases such as RSV pneumonia, cytomegalovirus (CMV) pneumonia, primary and metastatic lung cancer, and extranodal pulmonary non-Hodgkin's lymphoma; extrapulmonary diseases such as cancer and allograft rejection; and autoimmune diseases chosen from rheumatoid arthritis and Crohn's disease. The present sequence represents a fusion protein comprising the native human EPO signal peptide, human EPO and the human IgG1 Fc fragment (Fc-gamma-1), which is encoded by plasmid PED.dC.natEpoFc.		FT Region	Region /note= "Human mature EPO"
XX		FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"
OS Synthetic.		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
OS Chimeric.		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
XX		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
FR Key Location/Qualifiers		FT Region	Region /note= "Human mature EPO"
Peptide 1. 27		FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"
PT /label= EPO_signal_peptide		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
PT Protein 28. .428 "EPO/IgG1 Fc fragment fusion protein"		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
PT Region 28. .193		FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"
Query	Match 100.0%; Score 846; DB 8; Length 428;	FT Region	Region /note= "Human mature EPO"
Best Local Similarity 100.0%; Pred. No. 8. 9e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"	
Qy 1 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 60	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Db 28 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 87	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
SQ Sequence 428 AA;		FT Region	Region /note= "Human mature EPO"
Query	Match 100.0%; Score 846; DB 8; Length 428;	FT Region	Region /note= "8 residue peptide linker (SEQ ID NO:2)"
Best Local Similarity 100.0%; Pred. No. 8. 9e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Qy 1 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 60	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	
Db 28 APPRLICDSRVYERYLVEAKRAENTTGCABHCSLENINITYDPTKVNHYAKRMEVGQAA 87	FT Region	Region /note= "IgG1 Fc fragment_ (SEQ ID NO:2)"	

QY 121 PPDASAAAPLTTADTRKLFRVSNFLRGKUKLYGACRGD 165
 Db 148 PPDASAAAPLTTADTRKLFRVSNFLRGKUKLYGACRGD 192

RESULT 118
 ADW47520
 ID ADW47520 standard; protein; 435 AA.
 XX
 AC ADW47520;
 XX
 DT 24-MAR-2005 (first entry)
 XX
 DE Human EPO-linker-immunoglobulin Fc gamma 1 variant fusion protein.
 XX
 KW fusion protein; EPO; immunoglobulin.
 XX
 OS Homo sapiens.
 OS Synthetic.
 OS Unidentified.
 XX
 PN CN1521192-A.
 XX
 PD 18-AUG-2004.
 XX
 PP 30-JAN-2003; 2003CN-00115277.
 XX
 PR (XUHU-) XUHUA SHANGHAI BIOLOGY RES & DEV CO LTD.
 PA
 PT JIN Y, Sun N, Zhou R;
 XX
 DR WI; 2004-785669/78.
 DR N-PSDB; ADW47519.

XX
 PT Human erythropoietin Fc fusion protein with high bioactivity.
 XX
 PS Example 1; SEQ ID NO 22; 33pp; Chinese.

XX
 CC The invention relates to a novel human EPO and Fc fusion protein with similar or increased bioactivity to rHuEPO. The HuEPO-L-vFc fusion proteins of the invention contain human EPO, linked via a flexible peptide comprising 20 or less amino acids to a human IgG Fc variant, which has no lytic property and shows little Fc-mediating side effect. The invention further discloses the method for preparation of the fusion proteins. The HuEPO-L-vFc fusion protein may be useful for prolonging serum half-time, increasing bioactivity and improving the dynamic performance and effect of medicine. The current sequence is that of the human EPO-linker-immunoglobulin Fc gamma 1 variant fusion protein of the invention.

CC Sequence 435 AA;

SQ 100.0%; Score 846; DB 8; Length 435;
 Best Local Similarity 100.0%; Prede. No. 9.1e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERTLEAKEAENITGGARHCSLENNTTUPDKUNFYAKWKEVGQA 60
 Db 28 APPRLICDSRVLERTLEAKEAENITGGARHCSLENNTTUPDKUNFYAKWKEVGQA 87
 61 VEWVOGLALISSEAVRLQQLNLNSQPMELQLHQNDKAVSGLRSLTTLRAGQKEAIS 120
 Db 88 VEWVOGLALISSEAVRLQQLNLNSQPMELQLHQNDKAVSGLRSLTTLRAGQKEAIS 147

QY 121 PPDASAAAPLTTADTRKLFRVSNFLRGKUKLYGACRGD 165
 Db 148 PPDASAAAPLTTADTRKLFRVSNFLRGKUKLYGACRGD 192

RESULT 119
 AEA18937

ID	AE18937	
XX	standard; protein; 435 AA.	
AC	AE18937;	
XX		
DT	11-AUG-2005 (first entry)	
XX		
DE	Human erythropoietin-L-vFc-gamma1 fusion protein SEQ ID NO:22.	
XX		
KW	fusion protein; erythropoietin; IgG; immunoglobulin; immunotherapy; antianemic; anemia.	
XX		
OS	Homo sapiens.	
OS	Synthetic.	
XX		
FH	Key	Location/Qualifiers
FT	Peptide	1. -27
FT		/label= signal
FT	Protein	28.. 435
FT	Protein	/note= "HuEPO-L-vFc-gamma1 fusion protein"
FT	Protein	28.. 192
FT	Peptide	/note= "human erythropoietin amino acid sequence"
FT	Peptide	193.. 208
FT	Protein	/label= linker
FT	Protein	209.. 435
FT		/note= "Fc-gamma1 Ieu234Val, Ieu235Ala and Pro331Ser variant amino acid sequence"
XX		
PD	09-JUN-2005.	
XX		
PR	17-DEC-2004; 2004US-00016518.	
XX		
PR	17-AUG-2001; 2001US-00932812.	
XX		
PA	(SUNL/) SUN L K.	
PA	(SUNB/) SUN B N C.	
PA	(SUNC/) SUN C R Y.	
XX		
PI	Sun LK, Sun BNC, Sun CRY;	
XX		
DR	WPI; 2005-417006/42.	
XX		
PT	New recombinant HuEPO-L-vFc fusion protein Comprising HuEPO, a peptide linker, and a human IgG Fc variant, useful for treating anemia in patients caused by cancer chemotherapy, rheumatoid arthritis, myelodysplastic syndrome.	
PT	N-PSDB; AEA18936.	
XX		
CC	The invention relates to a recombinant HuEPO-L-vFc fusion protein consisting of human erythropoietin (HuEPO), a peptide linker, and a human IgG Fc variant, where the human IgG Fc variant comprises a hinge, CH ₂ , and CH ₃ domains of human IgG4 with Ser228Pro and Leu235Ala mutations as described: (1) a Chinese Hamster Ovary (CHO) cell line transfected with DNA encoding the recombinant HuEPO-L-vFc fusion protein in its growth period; and (2) a method for making the recombinant fusion protein comprising generating a CHO cell line transfected with DNA encoding the recombinant HuEPO-L-vFc fusion protein; growing the cell line under conditions the recombinant protein is expressed in its growth medium in excess of 10 microgram per million cells in a 24 hour period; and purifying the expressed protein, where the recombinant fusion protein exhibits an enhanced in vitro biological activity of at least 2 fold relative to that of rhEPO on a molar basis. The fusion protein is useful for treating anemia in patients caused by cancer chemotherapy, rheumatoid arthritis, arachniorine treatment for HIV infection and myelodysplastic syndrome. The HuEPO-L-vFc fusion proteins exhibit extended serum half-life and increases biological activities leading to improved pharmacokinetics and pharmacodynamics, and so fewer injections will be needed within a period of time. The present sequence represents the HuEPO-vFc-gamma1 fusion	

CC protein, which is used in the exemplification of the present invention.

XX DR WPI; 2005-457786/46.
XX N-PSDB; AER88756.

Query Match Score 846; DB 9; Length 435;
Best Local Similarity 100.0%; Pred. No. 9.1e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKENITGCAHCISLNENITVPDTKVNPFYAWKRMEMVGQQA 60
Db 28 APPRLICDSRVLERYLLEAKENITGCAHCISLNENITVPDTKVNPFYAWKRMEMVGQQA 87
Qy 61 VEWQGLALLSEAVLRGQALLVNSQWEPPLQLHDKAVSGLSLTILRALGAQKAIS 120
Db 88 VEWQGLALLSEAVLRGQALLVNSQWEPPLQLHDKAVSGLSLTILRALGAQKAIS 147
Qy 121 PPDASASAPLRTTADTPRKLFPRVYSNPLRGKLUKYGEACRTGD 165
Db 148 PPDASASAPLRTTADTPRKLFPRVYSNPLRGKLUKYGEACRTGD 192

SQ

RESULT 120

ID AEA8757
ID AEA8757 standard; protein; 435 AA.

XX AEA8757;

AC AEA8757;

DT 08-SEP-2005 (first entry)

DE Human erythropoletin (HuEPO)-L-vFc gamma fusion protein, SEQ ID: 22.

XX Fusion protein; erythropoletin; anemia; antianemic;
KW hematological disease; renal failure; nephrotropic;
KW genitorinary disease; rheumatoid arthritis; antiarthritic;
KW antineuramic; immune disorder; inflammation; musculoskeletal disease;
KW myelodysplastic syndrome; immunostimulant; neoplasm; IgG; antibody;
KW immunoglobulin; mutant; protein.

OS Homo sapiens.

OS Synthetic.

○

FH Key Location/Qualifiers
1. .27
FT /label= Signal_peptide

FT Protein
/note= "Mature human erythropoletin (HuEPO)-L-vFc gamma

FT fusion protein"

FT Region
28..192

/note= "Human erythropoletin (HuEPO)"

FT Region
193..208

/note= "Linker peptide"

FT Region
209..435

/note= "IgG variant (v) Fc gamma"

FT Misc-difference
222

/note= "Wild-type Leu substituted by Val"

FT Misc-difference
223

/note= "Wild-type Leu substituted by Ala"

FT /note= "Wild-type Pro substituted by Ser"

DN US2005142642-A1.

XX 30-JUN-2005.

PP 17-DEC-2004; 2004US-00017185.

PR 17-AUG-2001; 2001US-00932812.

XX (SUNL/) SUN L K.

PA (SUNB/) SUN B N C.

PA (SUNC/) SUN C R Y.

PI Sun LK, Sun BNC, Sun CRY;

XX DR WPI; 2005-457786/46.
XX N-PSDB; AER88756.

Query Match Score 846; DB 9; Length 435;
Best Local Similarity 100.0%; Pred. No. 9.1e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKENITGCAHCISLNENITVPDTKVNPFYAWKRMEMVGQQA 60
Db 28 APPRLICDSRVLERYLLEAKENITGCAHCISLNENITVPDTKVNPFYAWKRMEMVGQQA 87
Qy 61 VEWQGLALLSEAVLRGQALLVNSQWEPPLQLHDKAVSGLSLTILRALGAQKAIS 120
Db 88 VEWQGLALLSEAVLRGQALLVNSQWEPPLQLHDKAVSGLSLTILRALGAQKAIS 147
Qy 121 PPDASASAPLRTTADTPRKLFPRVYSNPLRGKLUKYGEACRTGD 165
Db 148 PPDASASAPLRTTADTPRKLFPRVYSNPLRGKLUKYGEACRTGD 192

SQ

RESULT 121

ID ADM33853
ID ADM33853 standard; protein; 436 AA.

XX ADM33853;

AC ADM33853;

DT 03-JUN-2004 (first entry)

DE Human HuEPO-L-vFc gamma 2 fusion protein.

XX Erythropoietin; EPO; immunoglobulin; IgG;
KW fragment crystallisation region; Fc; chronic anaemia; renal disease;
KW cancer chemotherapy; rheumatoid arthritis; AIDS;
KW myelodysplastic syndrome; (HuEPO)-L-vFc gamma 2; human.

OS Homo sapiens.

OS Synthetic.

○

FH Key Location/Qualifiers
1: .27
FT Peptide /note= "signal peptide"

FT Protein
/note= "EPO"

FT Peptide
/note= "Linker"

FT Protein
209..436

/note= "IgG2 Fc"

FT Misc-difference
390

/note= "Wild-type Pro substituted by Ser"

DN US2003082749-A1.

XX 01-MAY-2003.

XX	PR	17-AUG-2001; 2001US-00932812.	PP	antianaemic; nephrotropic; human; HuEPO-L-vFC; erythropoietin; EPO;
XX	PR	(SUNL/) SUN L K.	KW	anaemia; renal disease; cancer chemotherapy; rheumatoid arthritis;
PA	PA	(SUNB/) SUN B N C.	KW	AZT treatment; HIV infection; myelodysplastic syndrome; renal failure.
PA	PA	(SUNC/) SUN C R Y.	OS	Homo sapiens.
XX	PI	Sun LK, Sun BNC, sun CRY;	OS	Synthetic.
XX	XX	WPI; 2003-616080/5B.	XX	
DR	PT	New recombinant human erythropoietin-L-vFC fusion protein, useful for treating patients with chronic anaemia caused by renal failure, cancer chemotherapy, rheumatoid arthritis, or azathioprine treatment for HIV infection.	PT	
XX	PS	Claim 3; FIG 2A; 14pp; English.	PS	
XX	CC	The invention relates to a recombinant human erythropoietin (HuEPO) -L-vFC fusion protein comprising HuEPO, a peptide linker, and a human immunoglobulin G Fc (fragment crystallisation region) variant. Also included is a carbohydrate-derived cell line producing the human erythropoietin-L-vFC fusion protein cited above in its growth medium in excess of 10 microgramme per million cells in a 24-hour period. The HuEPO -L-vFC fusion protein exhibits an enhanced in vitro biological activity of at least 2-fold relative to that of recombinant HuEPO on a molar basis. The flexible peptide linker containing about 20 or fewer amino acids is present between HuEPO and the human IgG Fc variant. The IgG Fc contains amino acid mutations to attenuate effector functions. The human IgG Fc variant comprises a hinge, CH2 and CH3 domains of human IgG2 with Pro331Ser mutation, human IgG4 with Ser228Pro and Leu235Ala mutations, or human IgG1 with Leu235Ala and Pro331Ser mutations. The recombinant human erythropoietin-L-vFC fusion protein are useful for treating patients with chronic anaemia caused by renal failure, cancer chemotherapy, rheumatoid arthritis, azathioprine treatment for HIV infection, or myelodysplastic syndrome. The increased activity and prolonged presence of the human erythropoietin-L-vFC fusion protein in the serum, as compared to prior art, leads to lower dosages and less frequent injections. Less fluctuations of the drug in serum concentrations means improved safety and tolerability and less frequent injections result in better patient compliance and quality of life. The present sequence represents the fusion protein HuEPO-L-vFegamma2.	CC	
XX	CC	Sequence 436 AA;	CC	
XX	CC	Query Match Local Similarity 100.0%; Score 846; DB 7; Length 436; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;	CC	
QY	1	APRRLCDSRVLERVLRAKEAENTITGCACRHCSNLENITVDPDTKUNFYAWKRMENVGQDA 60	CC	
Dy	2B	APRRLCDSRVLERVLRAKEAENTITGCACRHCSNLENITVDPDTKUNFYAWKRMENVGQDA 87	CC	
QY	61	VEVMWGLALLSEAVIRGQALVNNSQPWERPLQLHDVKAVSGLRSITTLRALGAQEAIIS 120	CC	
Dy	88	VEVMWGLALLSEAVIRGQALVNNSQPWERPLQLHDVKAVSGLRSITTLRALGAQEAIIS 147	CC	
QY	121	PPDASASAPRTITATFERKLFRVSFNLFGKLYKGRCRTGD 165	CC	
Dy	148	PPDASASAPRTITATFERKLFRVSFNLFGKLYKGRCRTGD 192	CC	
RESULT	122		CC	
ADR48984			CC	
ID	ADR48984 standard; protein; 436 AA.		CC	
AC	ADR48984;		CC	
DT	02-DEC-2004 (first entry)		CC	
DE	HuEPO-L-Fc fusion protein.		CC	

CC protein of human erythropoietin (EPO), cDNA library of human fetal liver or kidney was used as the template in polymerase chain reaction (PCR). CC For the convenience of cloning, SEQ ID NO. 1 which incorporates a restriction enzyme cleavage site is used as the 5' oligonucleotide primer. The 3' primer (SEQ ID NO. 2) eliminates the EPO termination codon and incorporates a BamHI site. The resulting DNA fragments of approximately 600 bp were inserted into a holding vector such as pUC19 at the HindIII and BamHI sites to give the pBPO plasmid. The sequence of the CC human EPO gene was confirmed by DNA sequencing.

XX SQ Sequence 436 AA;

Query Match 100.0%; Score 846; DB 8; Length 436;
Best Local Similarity 100.0%; Pred. No. 9.1e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRKAENITTCGAECISLNENITVDPDKVNYAWKRMEVGQA 60
OY 28 APPRLICDSRVLERYLLEAKRKAENITTCGAECISLNENITVDPDKVNYAWKRMEVGQA 60

Db 61 VEWQGLALLSEAVTRGQLVNSQPEPLQLHVDKAVSGLSLTILRAKGKAIS 120
Qy 88 VEWQGLALLSEAVTRGQLVNSQPEPLQLHVDKAVSGLSLTILRAKGKAIS 120

Db 121 PPDASAPLRTTADPFKLFRVSYFLRGKLYGEACTGD 165
Qy 148 PPDASAPLRTTADPFKLFRVSYFLRGKLYGEACTGD 192

RESULT 123
ADW47516
ID ADW47516 standard; protein; 436 AA.
XX
AC
XX
DT 24-MAR-2005 (first entry)
XX Human EPO-linker-immunoglobulin Fc gamma 2 variant fusion protein.
XX fusion protein; EPO; immunoglobulin.
OS Synthetic.
XX Unidentified.
OS Homo sapiens.
XX
PR 30-JAN-2003; 2003CN-00115277.
XX
PA (XUHU-) XURHUA SHANGHAI BIOLOGY RES & DEV CO LTD.
XX
PI Jin Y, Sun N, Zhou R;
XX
DR WPI; 2004-785669/78.
XX
N-PSDB; ADW47515.
XX
PT Human erythropoietin Fc fusion protein with high bioactivity.
XX
PS Example 1; SEQ ID NO 18; 33pp; Chinese.

The invention relates to a novel human EPO and Fc fusion protein with proteins of the invention contain human EPO, linked via a flexible similar or increased bioactivity to HuEPO. The HuEPO-L-vFc fusion peptide comprising 20 or less amino acids, to a human IgG variant, which has no lytic property and shows little Fc-mediating side effect. The invention further discloses the method for preparation of the fusion protein. The HuEPO-L-vFc fusion protein may be useful for prolonging serum half-life, increasing bioactivity and improving the dynamic performance and effect of medicine. The current sequence is that of the

CC human EPO-linker-immunoglobulin Fc gamma 2 variant fusion protein of the invention.
CC XX SQ Sequence 436 AA;

Query Match 100.0%; Score 846; DB 8; Length 436;
Best Local Similarity 100.0%; Pred. No. 9.1e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRKAENITTCGAECISLNENITVDPDKVNYAWKRMEVGQA 60
OY 28 APPRLICDSRVLERYLLEAKRKAENITTCGAECISLNENITVDPDKVNYAWKRMEVGQA 60

Db 61 VEWQGLALLSEAVTRGQLVNSQPEPLQLHVDKAVSGLSLTILRAKGKAIS 120
Qy 88 VEWQGLALLSEAVTRGQLVNSQPEPLQLHVDKAVSGLSLTILRAKGKAIS 120

Db 121 PPDASAPLRTTADPFKLFRVSYFLRGKLYGEACTGD 165
Qy 148 PPDASAPLRTTADPFKLFRVSYFLRGKLYGEACTGD 192

RESULT 124
ID AEA18933
XX AEA18933 standard; protein; 436 AA.
AC AEA18933;
XX DT 11-AUG-2005 (first entry)
XX Human erythropoietin-L-vFc-gamma2 fusion protein SEQ ID NO:18.
XX fusion protein; erythropoietin; IgG; immunoglobulin; immunotherapy;
KW antianemic; anemia.
XX
OS Homo sapiens.
OS Synthetic.
XX
FH Key Location/Qualifiers
FT Peptide 1..27
FT /label= signal
FT Protein 28..436
FT /note= "HuEPO-L-vFc-gamma2 fusion protein"
FT Protein 28..436
FT /note= "human erythropoietin amino acid sequence"
FT Peptide 193..208
FT /label= linker
FT Protein 209..436
FT /note= "Fc-gamma2 Pro311Ser variant amino acid sequence"
XX
PN US2005124045-A1.
XX PD 09-JUN-2005.
XX
PF 17-DEC-2004; 2004US-00016518.
XX PR 17-AUG-2001; 2001US-00932812.
XX
PA (SUNL/) SUN L K.
PA (SUNB/) SUN B N C.
PA (SUNC/) SUN C R Y.
XX
PI Sun LK, Sun BNC, Sun CRY;
XX
DR WPI; 2005-417005/42.
N-PSDB; AEA18932.
XX
New recombinant HuEPO-L-vFc fusion protein comprising HuEPO, a peptide linker, and a human IgG Fc variant, useful for treating anemia in patients caused by cancer chemotherapy, rheumatoid arthritis, myelodysplastic syndrome.
XX Disclosure; SEQ ID NO 18; 24pp; English.

XX
 CC The invention relates to a recombinant HuEPO-L-vFc fusion protein
 CC consisting of human erythropoietin (HuEPO), a peptide linker, and a human
 CC IgG Fc variant, where the human IgG Fc variant comprises a hinge, CH2,
 CC and CH3 domains of human IgG4 with Ser228Pro and Leu235Ala mutations as
 CC described; (1) a Chinese Hamster Ovary (CHO) cell line transfected with
 CC DNA encoding the recombinant HuEPO-L-vFc fusion protein in its growth
 medium in excess of 10 or 30 micro gram per million cells in a 24 hour
 period; and (2) a method for making the recombinant fusion protein
 comprising generating a CHO cell line transfected with DNA encoding the
 recombinant HuEPO-L-vFc fusion protein, growing the cell line under
 conditions the recombinant protein is expressed in its growth medium in
 excess of 10 micro gram per million cells in a 24 hour period; and purifying
 the expressed protein, where the recombinant fusion protein exhibits an
 enhanced in vitro biological activity of at least 2 fold relative to that
 of rHuEPO on a molar basis. The fusion protein is useful for treating
 anemia in patients caused by cancer chemotherapy, rheumatoid arthritis,
 azathioprine treatment for HIV infection and myelodysplastic syndrome.
 The HuEPO-L-vFc fusion proteins exhibit extended serum half-life and
 increased biological activities, leading to improved pharmacokinetics and
 pharmacodynamics, and so fewer injections will be needed within a period
 of time. The present sequence represents the HuEPO-vFc-gamma2 fusion
 protein, which is used in the exemplification of the present invention.

XX
 CC Sequence 436 AA;

Query Match Score 846; DB 9; Length 436;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 60
 DB 28 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 87
 QY 61 VEWQGLALLSEAVLRGQALLNSSQPWFPLQLAVDKAWSGLSLSTTLRALGAQEAKS 120
 DB 88 VEWQGLALLSEAVLRGQALLNSSQPWFPLQLAVDKAWSGLSLSTTLRALGAQEAKS 147

QY 121 PPDASAPRLTTADTRKLFRVYSNPLRGKLYTGEACRGD 165

DB 148 PPDASAPRLTTADTRKLFRVYSNPLRGKLYTGEACRGD 192

Sequence 436 AA;

Query Match Score 846; DB 9; Length 436;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 60
 DB 28 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 87

DB Human erythropoietin (HuEPO)-L-vFc gamma2 fusion protein, SEQ ID: 18.

XX Fusion protein; erythropoietin; anemia; antianemic;
 XX hematological disease; nephrotropic;
 XX genitourinary disease; rheumatoid arthritis; antiarthritic;
 XX antirheumatic; immune disorder; inflammation; musculoskeletal disease;
 XX myelodysplastic syndrome; immunostimulant; neoplasm; IgG; antibody;
 XX immunoglobulin; muttein.
 OS Homo sapiens.
 OS Synthetic.

XX Key Location/Qualifiers
 PH 1. label= "Signal_peptide"
 FT 28 .436 /note= "Mature human erythropoietin (HuEPO)-L-vFc gamma2
 FT fusion protein"
 FT 28 .192 /note= "Human erythropoietin (HuEPO)"
 Region /note= "Human erythropoietin (HuEPO)"
 Region 193. .208

FT /note= "Linker peptide"
 FT Region /note= "Linker peptide"
 FT 209. .436 /note= "IgG variant (v) Fcgamma2"
 FT Msc-difference 320 /note= "Wild-type Pro substituted by Ser"
 FT /note= "Wild-type Pro substituted by Ser"
 XX PN US2005142642-A1.
 XX PD 30-JUN-2005.
 XX PP 17-DEC-2004; 2004US-00017185.
 PR 17-AUG-2001; 2001US-00932812.
 XX PA (SUNL/) SUN L K.
 PA (SUNB/) SUN B N C.
 PA (SUNC/) SUN C R Y.
 XX PT Sun LK, Sun BN, Sun CRV;
 DR WPI; 2005-457788/46.
 DR N-PSDB; AEA88752.
 PT New recombinant human erythropoietin (HuEPO)-L-vFc fusion protein, useful
 PT for managing anemia caused by conditions including renal failure, cancer
 PT chemotherapy, rheumatoid arthritis.
 XX Disclosure; SEQ ID NO 18; 24pp; English.
 CC The present invention relates to a recombinant human erythropoietin
 CC (HuEPO)-L-variant (v) Fc fusion protein comprising HuEPO, a peptide
 CC linker and a human immunoglobulin G (IgG) Fc variant, where the human IgG
 CC Fc variant comprises a hinge, CH2 and CH3 domains; human IgG1 with
 CC Leu235Ala, Leu255Ala and Pro335Ser mutations. The recombinant protein is
 CC useful for treating anemia caused by conditions including renal failure,
 CC infection and myelodysplastic syndrome. The present sequence is a HuEPO-L
 CC -vFc gamma2 fusion protein.
 XX Sequence 436 AA;

Query Match Score 846; DB 9; Length 436;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 60
 DB 28 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 87

DB Human erythropoietin (HuEPO)-L-vFc gamma4 fusion protein, SEQ ID: 18.

XX Fusion protein; erythropoietin; anemia; antianemic;
 XX hematological disease; nephrotropic;
 XX genitourinary disease; rheumatoid arthritis; antiarthritic;
 XX antirheumatic; immune disorder; inflammation; musculoskeletal disease;
 XX myelodysplastic syndrome; immunostimulant; neoplasm; IgG; antibody;
 XX immunoglobulin; muttein.
 OS Homo sapiens.
 OS Synthetic.

XX Key Location/Qualifiers
 PH 1. label= "Signal_peptide"
 FT 28 .436 /note= "Mature human erythropoietin (HuEPO)-L-vFc gamma2
 FT fusion protein"
 FT 28 .192 /note= "Human erythropoietin (HuEPO)"
 Region /note= "Human erythropoietin (HuEPO)"
 Region 193. .208

FT /note= "Linker peptide"
 FT Region /note= "Linker peptide"
 FT 209. .436 /note= "IgG variant (v) Fcgamma2"
 FT Msc-difference 320 /note= "Wild-type Pro substituted by Ser"
 FT /note= "Wild-type Pro substituted by Ser"
 XX PN US2005142642-A1.
 XX PD 30-JUN-2005.
 XX PP 17-DEC-2004; 2004US-00017185.
 PR 17-AUG-2001; 2001US-00932812.
 XX PA (SUNL/) SUN L K.
 PA (SUNB/) SUN B N C.
 PA (SUNC/) SUN C R Y.
 XX PT Sun LK, Sun BN, Sun CRV;
 DR WPI; 2005-457788/46.
 DR N-PSDB; AEA88752.
 PT New recombinant human erythropoietin (HuEPO)-L-vFc fusion protein, useful
 PT for managing anemia caused by conditions including renal failure, cancer
 PT chemotherapy, rheumatoid arthritis.
 XX Disclosure; SEQ ID NO 18; 24pp; English.
 CC The present invention relates to a recombinant human erythropoietin
 CC (HuEPO)-L-variant (v) Fc fusion protein comprising HuEPO, a peptide
 CC linker and a human immunoglobulin G (IgG) Fc variant, where the human IgG
 CC Fc variant comprises a hinge, CH2 and CH3 domains; human IgG1 with
 CC Leu235Ala, Leu255Ala and Pro335Ser mutations. The recombinant protein is
 CC useful for treating anemia caused by conditions including renal failure,
 CC infection and myelodysplastic syndrome. The present sequence is a HuEPO-L
 CC -vFc gamma2 fusion protein.
 XX Sequence 436 AA;

Query Match Score 846; DB 9; Length 436;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 60
 DB 28 APPRLICDSRVLERYLLEAKRAENITTCGAECRHSCLNENITVPTDKVNFKVAKWKGMEVGQOA 87

DB Human erythropoietin (HuEPO)-L-vFc gamma4 fusion protein, SEQ ID: 18.

XX Fusion protein; erythropoietin; anemia; antianemic;
 XX hematological disease; nephrotropic;
 XX genitourinary disease; rheumatoid arthritis; antiarthritic;
 XX antirheumatic; immune disorder; inflammation; musculoskeletal disease;
 XX myelodysplastic syndrome; immunostimulant; neoplasm; IgG; antibody;
 XX immunoglobulin; muttein.
 OS Homo sapiens.
 OS Synthetic.

XX Key Location/Qualifiers
 PH 1. label= "Signal_peptide"
 FT 28 .436 /note= "Mature human erythropoietin (HuEPO)-L-vFc gamma2
 FT fusion protein"
 FT 28 .192 /note= "Human erythropoietin (HuEPO)"
 Region /note= "Human erythropoietin (HuEPO)"
 Region 193. .208

OS Homo sapiens.
 OS Synthetic.
 XX
 FT Key
 FT Peptide
 FT Protein
 FT Peptide
 FT Peptide
 FT Protein
 FT Misc-difference
 FT Misc-difference
 FT Misc-difference
 FT PN US2003082749-A1.
 XX PD 01-MAY-2003.
 XX PP 17-AUG-2001; 2001US-00932812.
 XX PR 17-AUG-2001; 2001US-00932812.
 XX PA (SUNL/) SUN L K.
 PA (SUNB/) SUN B N C.
 PA (SUNC/) SUN C R Y.
 XX PI Sun LK, Sun BNC, Sun CRY;
 XX DR WPI; 2003-6116080/58.
 XX N-PSB; APM33854.
 XX PT New recombinant human erythropoietin-L-vFC fusion proteins, useful for treating patients with chronic anemia caused by renal failure, cancer chemotherapy, rheumatoid arthritis, or azathioprine treatment, for HIV infection.
 XX PT New recombinant human erythropoietin-L-vFC fusion protein comprising a hinge, CH₂ and CH₃ domains of human IgG2 with IGG Fc variant comprises a hinge, CH₂ and CH₃ domains of human IgG4 with Ser28Pro and Leu235Ala mutations, or human IgG1 with Leu234Val, Leu235Ala and Pro31Ser mutations. The recombinant human erythropoietin-L-vFC fusion proteins are useful for treating patients with chronic anemia caused by renal failure, cancer chemotherapy, rheumatoid arthritis, azathioprine treatment for HIV infection, or myelodysplastic syndrome. The increased activity and prolonged presence of the human erythropoietin-L-vFC fusion protein in the serum, as compared to prior art, leads to lower dosages and less frequent injections. Less fluctuations of the drug in serum concentrations means improved safety and tolerability, and less frequent injections result in better patient compliance and quality of life. The present sequence represents the fusion protein HuEPO-L-vFCgamma4.
 XX SQ Sequence 437 AA;

The invention relates to a recombinant human erythropoietin (HuEPO) -L-vFC fusion protein comprising HuEPO, a peptide linker, and a human immunoglobulin G Fc (fragment crystallization region) variant. Also included is a carbohydrate-derived cell line producing the human erythropoietin-L-vFC fusion protein cited above in its growth medium in excess of 10 microgramme per million cells in a 24-hour period. The HuEPO -L-vFC fusion protein exhibits an enhanced in vitro biological activity of at least 2-fold relative to that of recombinant HuEPO on a molar basis. The flexible peptide linker containing about 20 or fewer amino acids is present between HuEPO and the human IgG Fc variant. The IgG Fc contains amino acid mutation to attenuate effector functions. The human IgG Fc variant comprises a hinge, CH₂ and CH₃ domains of human IgG2 with Pro31Ser mutation, human IgG4 with Ser28Pro and Leu235Ala mutations, or human IgG1 with Leu234Val, Leu235Ala and Pro31Ser mutations. The recombinant human erythropoietin-L-vFC fusion proteins are useful for treating patients with chronic anemia caused by renal failure, cancer chemotherapy, rheumatoid arthritis, azathioprine treatment for HIV infection, or myelodysplastic syndrome. The increased activity and prolonged presence of the human erythropoietin-L-vFC fusion protein in the serum, as compared to prior art, leads to lower dosages and less frequent injections. Less fluctuations of the drug in serum concentrations means improved safety and tolerability, and less frequent injections result in better patient compliance and quality of life. The present sequence represents the fusion protein HuEPO-L-vFCgamma4.

XX SQ Sequence 437 AA;

Query Match 100.0%; Score 846; DB 7; Length 437;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
 CC 28 APPRLICDSVRLVILEAKAENITTGCAEHCSLNENITPDTKQFYAWKRMEVGQQA 87
 CC 61 VEWVGQALISEAVIRGQALLVNSQQPWEFLQLRHDKAISGLRSLTLLRAGQKAIS 120
 CC 88 VEWVGQALISEAVIRGQALLVNSQQPWEFLQLRHDKAISGLRSLTLLRAGQKAIS 147
 CC 121 PPDAAASAPLRTTADTFRKLFRVYSNFLRGKLKLYTGACRTGD 165
 CC 148 PPDAAASAPLRTTADTFRKLFRVISNFLRGKLKLYTGACRTGD 192

Db RESULT 127
 DR ADR48986
 ID ADR48986 standard; protein; 437 AA.
 XX AC ADR48986;
 XX DT 02-DEC-2004 (first entry)
 XX DR HuEPO-L-vFc fusion protein #1.
 XX KW antianemic; nephrotropic; human; HuEPO-L-vFc; erythropoietin; EPO;
 KW anaemia; renal disease; cancer chemotherapy; rheumatoid arthritis;
 KW AZT treatment; HIV infection; myelodysplastic syndrome; renal failure.
 XX PR Homo sapiens.
 XX OS Synthetic.
 XX PN US2004175824-A1.
 XX DR 09-SEP-2004.
 XX PP 21-JAN-2004; 2004US-00761593.
 XX PR 17-AUG-2001; 2001US-00932812.
 XX PA (SUNL/) SUN L K.
 PA (SUNB/) SUN B N C.
 PA (SUNC/) SUN C R Y.
 XX PI Sun LK, Sun BNC, Sun CRY;
 XX DR WPI; 2004-634851/61.
 XX N-PSB; ADR48985.
 XX PT New recombinant HuEPO-L-vFC fusion protein comprises human erythropoietin (HuEPO), a peptide linker, and a human IgG Fc variant, useful for treating chronic anemia due to renal diseases, cancer chemotherapy, or rheumatoid arthritis.
 XX PR Claim 4; SEQ ID NO 20; 31pp; English.
 XX PS A recombinant HuEPO-L-vFC fusion protein comprises human erythropoietin (HuEPO), a peptide linker, and a human IgG Fc variant. is new. INDEPENDENT CLAIMS are also included for the following: a chinese hamster ovary (CHO)-derived cell line producing the HuEPO-L-vFc fusion protein in its growth medium in excess of 10 microg per million cells in a 24 hour period; and a method for making a recombinant fusion protein comprising HuEPO, a flexible peptide linker, and a human IgG Fc variant. Preferred protein: The peptide linker containing 20 or fewer amino acids is present between HuEPO and the human IgG Fc variant, and comprises two or more amino acids selected from glycine, serine, alanine, and threonine. The human IgG Fc variant comprises a hinge, CH₂, and CH₃ domains of human IgG2 with Pro31Ser mutation comprising 436 amino acids (SEQ ID NO. 18). It also comprises a hinge, CH₂, and CH₃ domains of human IgG4 with Ser28Pro and Leu235Ala mutations comprising 437 amino acids (SEQ ID NO. 20). It further comprises a hinge, CH₂, and CH₃ domains of human IgB1 with Leu234Val, Leu235Ala, and Pro31Ser mutations comprising 435 amino acids (SEQ ID NO. 22). The HuEPO-L-vFC fusion protein exhibits in vitro biological activity similar to or higher than that of HuEPO on a molar basis. Preferred CHO-Derived Cell line: The CHO-derived cell line producing the HuEPO-L-vFC fusion protein in its growth medium in excess

CC of 30 µg per million cells in a 24 hour period. The human IgG FC variant comprises a hinge, CH2, CH3 domains of human IgG selected from Igb1 as SEQ ID NO. 22, IgG2 as SEQ ID NO. 18, and IgG4 as SEQ ID NO. 20, the IgG FC contains amino acid mutations to attenuate effector functions, a flexible peptide linker containing 20 or fewer amino acids is present between HuRPO and human IgG FC variant, and the HuRPO-L-vFc fusion protein exhibits in vitro biological activity similar to or higher than that of rhRPO on a molar basis. Preferred Method: Making a recombinant fusion protein comprising HuRPO, a flexible peptide linker, and a human IgG FC variant comprises: generating a CHO-derived cell line, growing the cell line where the recombinant protein is expressed in its growth medium in excess of 10 µg per million cells in a 24 hour period; and purifying the expressed protein from (b), where the recombinant fusion protein exhibits in vitro biological activity similar to or higher than that of rhRPO on a molar basis. Antianemic; Nephrotropic. No biological data given. None given. Administration can be through subcutaneous or intravenous route. No dosage given. The recombinant HuRPO-L-vFc fusion protein is useful for treating patients with chronic anemia due to renal diseases, cancer chemotherapy, rheumatoid arthritis, AZT treatment for HIV infection, or myelodysplastic syndrome. It is also useful in the treatment of renal failure. A fusion protein was assembled from several DNA segments. To obtain the gene encoding the leader peptide and mature protein of human erythropoietin (EPO), cDNA library of human fetal liver or kidney was used as the template in polymerase chain reaction (PCR). For the convenience of cloning, SEQ ID NO. 1 which incorporates a restriction enzyme cleavage site is used as the 5' oligonucleotide primer. The 3' primer (SEQ ID NO. 2) eliminates the EPO termination codon and incorporates a BamHI site. The resulting DNA fragments of approximately 600 bp were inserted into a holding vector such as pUC19 at the HindIII and BamHI sites to give the pEPO plasmid. The sequence of the human EPO gene was confirmed by DNA sequencing.

XX SQ Sequence 437 AA;

Query Match 100.0%; Score 846; DB 8; Length 437;
Best Local Similarity 100.0%; Pred. No. 9.1e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 APPRLICDSRVLRVYLRAKEAENITTGCAEHCSLNENITVPTKVNFTYAWKRMEVQQAA 60
Db 28 APPRLICDSRVLRVYLRAKEAENITTGCAEHCSLNENITVPTKVNFTYAWKRMEVQQAA 87

OY 61 VEWQGLALLSEAVLQRQALLYNSQPWEPLQLHVDKAVSGLSLTTLRALGAQEAIIS 120
Db 88 VEWQGLALLSEAVLQRQALLYNSQPWEPLQLHVDKAVSGLSLTTLRALGAQEAIIS 147

RESULT 128
ID ADW47518 standard; protein; 437 AA.
AC AC
XX
DT 24-MAR-2005 (first entry)

DE Human EPO-linker-immunoglobulin Fc gamma 4 variant fusion protein.
KW fusion protein; EPO; immunoglobulin.
OS Homo sapiens.
OS Synthetic.
OS Unidentified.
XX CN1521192-A.
PN 18-AUG-2004.
XX 30-JAN-2003; 2003CN-00115277.
XX

PR 30-JAN-2003; 2003CN-00115277.
XX (XUHU-) XUTUA SHANGHAI BIOLOGY RES & DEV CO LTD.
PA
XX PI Jin Y, Sun N, Zhou R;
XX DR WPI; 2004-785669/78.
N-PSDB; ADW47517.

XX PR Human erythropoietin Fc fusion protein with high bioactivity.
XX RS Example 1; SEQ ID NO 20; 33PP; Chinese.

XX SQ Sequence 437 AA;
Query Match 100.0%; Score 846; DB 8; Length 437;
Best Local Similarity 100.0%; Pred. No. 9.1e-86;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

OY 1 APPRLICDSRVLRVYLRAKEAENITTGCAEHCSLNENITVPTKVNFTYAWKRMEVQQAA 60
Db 28 APPRLICDSRVLRVYLRAKEAENITTGCAEHCSLNENITVPTKVNFTYAWKRMEVQQAA 87

OY 61 VEWQGLALLSEAVLQRQALLYNSQPWEPLQLHVDKAVSGLSLTTLRALGAQEAIIS 120
Db 88 VEWQGLALLSEAVLQRQALLYNSQPWEPLQLHVDKAVSGLSLTTLRALGAQEAIIS 147

RESULT 129
ID AEA18935 standard; protein; 437 AA.
AC AEA18935;
XX
DT 11-AUG-2005 (first entry)

DE Human erythropoietin-L-vFc-gamma4 fusion protein SEQ ID NO:20.
XX KW fusion protein; erythropoietin; IgG; immunoglobulin; immunotherapy;
XX OS Homo sapiens.
OS Synthetic.

XX PH Key Location/Qualifiers
PT Peptide 1..27
PT /label= signal
PT Protein 28..437
PT /note= "HuRPO-L-vFc-gamma4 fusion protein"
PT Protein 28..192
PT /note= "human erythropoietin amino acid sequence."
PT Peptide 193..208
PT /label= linker
PT Protein 209..437
PT /note= "Fc-gamma4 Ser228Pro and Leu235Ala variant amino acid sequence".
XX PF PF PF PF

PN US2005124045-A1.
 XX DT 08-SEP-2005 (first entry)
 PD XX Human erythropoietin (HuEPO)-L-vFcgamma4 fusion protein, SEQ ID: 20.
 KK DE
 KK Fusion protein; erythropoietin; anemia; antianemic;
 KW hematochemical disease; renal failure; nephrotropic;
 KW genitourinary disease; rheumatoid arthritis; antiarthritic;
 KW anti-rheumatic; immune disorder; inflammation; musculoskeletal disease;
 KW myelodysplastic syndrome; immunostimulant; neoplasm; IgG; antibody;
 KW immunoglobulin; mutein.
 XX OS Homo sapiens.
 OS Synthetic.

XX Key Location/Qualifiers
 PR Peptide 1..27
 FT /label= Signal_peptide
 FT Protein 28..437
 /note= "Mature human erythropoietin (HuEPO)-L-vFcgamma4 fusion protein"
 FT Region 28..192
 /note= "Human erythropoietin (HuEPO)"
 FT Region 193..208
 /note= "Linker peptide"
 FT Region 209..437
 /note= "IgG variant (v) Fcgamma4"
 FT Misc-difference 218
 /note= "Wild-type Ser substituted by Pro"
 FT Misc-difference 225
 /note= "Wild-type Leu substituted by Ala"
 XX PR 17-AUG-2001; 2001US-00932812.
 XX PN US2005142642-A1.
 XX PD 30-JUN-2005.
 XX PF 17-DEC-2004; 2004US-00017185.
 XX PR 17-AUG-2001; 2001US-00932812.
 XX PA (SUNL/) SUN L K.
 PA (SUNB/) SUN B N C.
 PA (SUNC/) SUN C R Y.
 XX PI Sun LK, Sun BNC, Sun CRY;
 XX DR WPI; 2005-457788/46.
 DR N-PSDB; AEA88754.
 XX PR New recombinant human erythropoietin (HuEPO)-L-vFc fusion protein, useful
 PT for managing anemia caused by conditions including renal failure, cancer
 PT chemotherapy, rheumatoid arthritis.
 XX PS Disclosure; SEQ ID NO 20; 24pp; English.
 XX The present invention relates to a recombinant human erythropoietin
 (rHuEPO)-L-variant (v) FC fusion protein comprising HuEPO, a peptide
 linker and a human immunoglobulin G (IgG) FC variant, where the human IgG
 FC Variant comprises a hinge, CH2 and CH3 domains of human IgG1 with
 Leu235Val, Leu235Ala and Pro331Ser mutations. The recombinant protein is
 useful for treating anemia caused by conditions including renal failure,
 cancer chemotherapy, rheumatoid arthritis, AZT treatment for HIV
 infection and myelodysplastic syndrome. The present sequence is a HuEPO-L
 -vFc gamma4 fusion protein.
 XX SQ Sequence 437 AA;

Query Match 100 %; Score 846; DB 9; Length 437;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTCGAECHSLNENITVPTDKVNFYAWKRMVGQQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITTCGAECHSLNENITVPTDKVNFYAWKRMVGQQA 87
 QY 61 VEWQGLALLSEAVLRGQALLNSQPEPLQLAVDRAVSGLRSLTILRAAGKAERAS 120
 Db 88 VEWQGLALLSEAVLRGQALLNSQPEPLQLAVDRAVSGLRSLTILRAAGKAERAS 147
 QY 121 PPDASASAPLTTTADTPFRKLFRVYSFLRGKLUKYGEACRTGD 165
 Db 148 PPDASASAPLTTTADTPFRKLFRVYSFLRGKLUKYGEACRTGD 192

XX SQ Sequence 437 AA;

Query Match 100 %; Score 846; DB 9; Length 437;
 Best Local Similarity 100.0%; Pred. No. 9.1e-86; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTCGAECHSLNENITVPTDKVNFYAWKRMVGQQA 60
 Db 28 APPRLICDSRVLERYLLEAKEAENITTCGAECHSLNENITVPTDKVNFYAWKRMVGQQA 87

QY 61 VEWQGLLISEAVLRGQALVNSQWPFLQLAVDKAVGSLSLTIRALGAQKAEIS 120
 CC ||||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
 CC 88 VEWQGLLISEAVLRGQALVNSQWPFLQLAVDKAVGSLSLTIRALGAQKAEIS 147
 CC ||||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
 QY 121 PPDRASAAPLRTITADTPRKLFRLFRVYSNFLRGKLUKYGEACRGD 165
 CC ||||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
 Db 148 PPDRASAAPLRTITADTPRKLFRLFRVYSNFLRGKLUKYGEACRGD 192
 CC ||||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||

RESULT 131

ADP1655

ID ADP1655 standard; protein; 768 AA.

XX AC ADP1655;

XX DT 12-FEB-2004 (first entry)

Human albumin therapeutic fusion protein SeqID162.

KW albumin fusion protein; albumin activity; human serum albumin;
 serum osmotic pressure; shelf-life; stability; antidiabetic;
 gene therapy; diabetes mellitus; human.

XX OS Chimeric.

OS Homo sapiens.

XX PN WO2003060071-A2.

XX PD 24-JUL-2003.

23-DEC-2002; 2002WO-US040891.

XX PR 21-DEC-2001; 2001US-0341811P.

XX PR 24-JAN-2002; 2002US-0340358P.

XX PR 28-JAN-2002; 2002US-0351360P.

PR 26-FEB-2002; 2002US-0359370P.

PR 28-FEB-2002; 2002US-0360000P.

PR 27-MAR-2002; 2002US-0367500P.

PR 08-APR-2002; 2002US-0370227P.

PR 10-MAY-2002; 2002US-0378950P.

PR 24-MAY-2002; 2002US-0382617P.

PR 28-MAY-2002; 2002US-0383123P.

PR 05-JUN-2002; 2002US-0385708P.

PR 10-JUL-2002; 2002US-03864625P.

PR 24-JUL-2002; 2002US-0388008P.

PR 09-AUG-2002; 2002US-0402131P.

PR 13-AUG-2002; 2002US-0402708P.

PR 18-SEP-2002; 2002US-0411355P.

PR 02-OCT-2002; 2002US-0414984P.

PR 11-OCT-2002; 2002US-0417611P.

PR 23-OCT-2002; 2002US-0420246P.

PR 05-NOV-2002; 2002US-0423623P.

XX PA (HUMA-) HUMAN GENOME SCI INC.

PA (DELB-) DELTA BIOTECHNOLOGY LTD.

PA (PRIN-) PRINCIPIA PHARM CORP.

XX PI Ballance DJ, Turner AJ, Roben CA, Haseltine WA;

XX DR WPI; 2003-598517/56.

XX XX New albumin fusion protein, useful for preparing a composition for

PT treating diabetes mellitus.

XX PS Example 4; SEQ ID NO 162; 24pp; English.

the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is the amino acid sequence of a novel full-length human albumin therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at [ftp.wipo.int/pub/publishedpat_sequences](http://wipo.int/pub/publishedpat_sequences)

Sequence 768 AA;

Query Match 100.0%; score 846; DB 7; Length 768;
 Best Local Similarity 100.0%; Pred. No. 2.1e-85; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Delins 0;

QY 1 APPLICDSRVLEVLRLAKAENITTCGAECIHSCLNNNTIVDTKVNPFYAWKGRMEVQQA 60
 Db 604 APPRLICDSRVLEVLRLAKAENITTCGAECIHSCLNNNTIVDTKVNPFYAWKGRMEVQQA 663
 QY 61 VEWQGLLISEAVLRGQALVNSQWPFLQLAVDCAVSGRSLSLTIRALGAQKAEIS 120
 Db 664 VEWQGLLISEAVLRGQALVNSQWPFLQLAVDCAVSGRSLSLTIRALGAQKAEIS 723

QY 121 PPDRASAAPLRTITADTPRKLFRLFRVYSNFLRGKLUKYGEACRGD 165
 Db 724 PPDRASAAPLRTITADTPRKLFRLFRVYSNFLRGKLUKYGEACRGD 768

RESULT 132

ADP16425

ID ADP16425 standard; protein; 768 AA.

XX AC ADP16425;

XX DT 12-FEB-2004 (first entry)

XX DE Human albumin therapeutic fusion protein SeqID1522.

KW albumin fusion protein; albumin activity; human serum albumin;

serum osmotic pressure; shelf-life; stability; antidiabetic;

gene therapy; diabetes mellitus; human.

XX OS Chimeric.

OS Homo sapiens.

XX PN WO2003060071-A2.

XX PD 24-JUL-2003.

XX PR 23-DEC-2002; 2002WO-US040891.

XX PR 21-DEC-2001; 2001US-0341811P.

PR 24-JAN-2002; 2002US-0350350P.

PR 28-JAN-2002; 2002US-0351360P.

PR 26-FEB-2002; 2002US-0359370P.

PR 28-FEB-2002; 2002US-0360000P.

PR 27-MAR-2002; 2002US-0367500P.

PR 24-JUL-2002; 2002US-0370227P.

PR 08-APR-2002; 2002US-0378950P.

PR 10-MAY-2002; 2002US-0382617P.

PR 24-MAY-2002; 2002US-0383123P.

PR 28-MAY-2002; 2002US-0411355P.

PR 05-JUN-2002; 2002US-0385708P.

PR 10-JUL-2002; 2002US-0394675P.

PR 23-OCT-2002; 2002US-0417611P.

PR 23-OCT-2002; 2002US-0420246P.

This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of

XX (HUMA-) HUMAN GENOME SCI INC.
 PA (DELBZ) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPIA PHARM CORP.
 XX
 PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX DR WPI; 2003-598517/56.
 PT New albumin fusion protein, useful for preparing a composition for
 PT treating diabetes mellitus.
 XX PS Example 4; SEQ ID NO 1522; 24pp; English.
 XX This invention relates to a novel albumin fusion protein having albumin
 CC or biological activity. Human serum albumin is responsible for a
 CC significant proportion of the osmotic pressure of serum and also
 CC functions as a carrier of endogenous and exogenous ligands. The fusion of
 CC albumin to a therapeutic protein may increase shelf-life and stability of
 CC the therapeutic protein. The albumin fusion protein of the invention may
 CC allow production of compositions with antidiabetic activity whilst the
 CC nucleotide sequence which encodes it may be useful for gene therapy. The
 CC albumin fusion protein is useful for preparing a composition for treating
 CC diabetes mellitus. The present sequence is the amino acid sequence of a
 CC novel full-length human albumin therapeutic fusion protein of the
 CC invention. Note: The sequence data for this patent did not form part of
 CC the printed specification, but was obtained in electronic format directly
 CC from WIPO at ftp://ftp.wipo.int/pub/publishedpat_sequences
 XX SQ Sequence 768 AA;

Query Match 100.0%; Score 846; DB 7; Length 768;
 Best Local Similarity 100.0%; Pred. No. 2.1e-85; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 APPRLICDSRVIERYLLEAKEAEINTTGCABHCSLNENITVDPDKVNFYAWKRMEMVGQAA 60
 Db 604 APPRLICDSRVIERYLLEAKEAEINTTGCABHCSLNENITVDPDKVNFYAWKRMEMVGQAA 663
 Oy 61 VEWVQGALLSEAVLRGQALLVNSSQPWEPLQLHDKAWSGLRSLTLLRAIGAQEAIS 120
 Db 664 VEWVQGALLSEAVLRGQALLVNSSQPWEPLQLHDKAWSGLRSLTLLRAIGAQEAIS 723
 Oy 121 PPDAASAPLRTTADTPRKLFRVVSNFLRGKLUYTGEACRTGD 165
 Db 724 PPDAASAPLRTTADTPRKLFRVVSNFLRGKLUYTGEACRTGD 768

XX PT New albumin fusion protein, useful for preparing a composition for
 XX treating diabetes mellitus.
 XX PS Example 4; SEQ ID NO 1661; 24pp; English.
 XX This invention relates to a novel albumin fusion protein having albumin
 CC or biological activity. Human serum albumin is responsible for a
 CC significant proportion of the osmotic pressure of serum and also
 CC functions as a carrier of endogenous and exogenous ligands. The fusion of
 CC albumin to a therapeutic protein may increase shelf-life and stability of
 CC the therapeutic protein. The albumin fusion protein of the invention may
 CC allow production of compositions with antidiabetic activity whilst the
 CC nucleotide sequence which encodes it may be useful for gene therapy. The
 CC albumin fusion protein is useful for preparing a composition for treating
 CC diabetes mellitus. The present sequence is the amino acid sequence of a
 CC novel full-length human albumin therapeutic fusion protein of the
 CC invention. Note: The sequence data for this patent did not form part of
 CC the printed specification, but was obtained in electronic format directly
 CC from WIPO at ftp://ftp.wipo.int/pub/publishedpat_sequences
 XX SQ Sequence 768 AA;

Query Match 100.0%; Score 846; DB 7; Length 768;
 Best Local Similarity 100.0%; Pred. No. 2.1e-85; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Oy 1 APPRLICDSRVIERYLLEAKEAEINTTGCABHCSLNENITVDPDKVNFYAWKRMEMVGQAA 60
 Db 604 APPRLICDSRVIERYLLEAKEAEINTTGCABHCSLNENITVDPDKVNFYAWKRMEMVGQAA 663
 Oy 61 VEWVQGALLSEAVLRGQALLVNSSQPWEPLQLHDKAWSGLRSLTLLRAIGAQEAIS 120
 Db 664 VEWVQGALLSEAVLRGQALLVNSSQPWEPLQLHDKAWSGLRSLTLLRAIGAQEAIS 723
 Oy 121 PPDAASAPLRTTADTPRKLFRVVSNFLRGKLUYTGEACRTGD 165
 Db 724 PPDAASAPLRTTADTPRKLFRVVSNFLRGKLUYTGEACRTGD 768

XX RESULT 133
 ADF16564
 ID ADF16564 standard; protein; 768 AA.
 XX AC ADF16564;
 XX DT 12-FEB-2004 (first entry)
 DE Human albumin therapeutic fusion protein SeqId1661.
 XX KW albumin fusion protein; albumin activity; human serum albumin;
 KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW gene therapy; diabetes mellitus; human.
 OS Chimeric.
 OS Homo sapiens.
 PN WO2003060071-A2.
 XX PD 24-JUL-2003.
 XX ID ADF16426
 XX ID ADF16426 standard; protein; 768 AA.
 XX AC ADF16426;
 XX PR 21-DEC-2001; 2001US-0341811P.
 XX PR 24-JAN-2002; 2002US-0351360P.
 PR 26-FEB-2002; 2002US-0359370P.
 PR 28-FEB-2002; 2002US-0360000P.
 PR 27-MAR-2002; 2002US-0367500P.
 PR 08-APR-2002; 2002US-0370227P.
 PR 10-MAY-2002; 2002US-037950P.
 PR 24-MAY-2002; 2002US-0382017P.
 PR 28-MAY-2002; 2002US-0383123P.
 PR 05-JUN-2002; 2002US-0385708P.
 PR 10-JUN-2002; 2002US-0394625P.
 PR 24-JUN-2002; 2002US-0398008P.
 PR 09-AUG-2002; 2002US-0402131P.
 PR 13-AUG-2002; 2002US-0402708P.
 PR 18-SEP-2002; 2002US-0411355P.
 PR 18-SEP-2002; 2002US-0411426P.
 PR 02-OCT-2002; 2002US-0414984P.
 PR 11-OCT-2002; 2002US-0417611P.
 PR 23-OCT-2002; 2002US-0420246P.
 PR 05-NOV-2002; 2002US-0423623P.
 XX (HUMA-) HUMAN GENOME SCI INC.
 PA (DELBZ) DELTA BIOTECHNOLOGY LTD.
 PA (PRIN-) PRINCIPIA PHARM CORP.
 XX PT Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX DR WPI; 2003-598517/56.

DT DT 12-FEB-2004 (first entry)
 XX XX Human albumin therapeutic fusion protein seqID1523.
 DB DB albumin fusion protein; albumin activity; human serum albumin;
 KW KW gene therapy; diabetes mellitus; human.
 XX XX Chimeric.
 OS OS Homo sapiens.
 XX XX WO2003060071-A2.
 PD PD 24-JUL-2003.
 PP PP 21-DEC-2002; 2002WO-US040891.
 PR PR 21-DEC-2001; 2001US-0341811P.
 PR PR 24-JAN-2002; 2002US-035038P.
 PR PR 28-JAN-2002; 2002US-0311360P.
 PR PR 28-FEB-2002; 2002US-0359370P.
 PR PR 21-MAR-2002; 2002US-0367500P.
 PR PR 01-APR-2002; 2002US-0370227P.
 PR PR 10-MAY-2002; 2002US-0378950P.
 PR PR 24-MAY-2002; 2002US-0382617P.
 PR PR 05-JUN-2002; 2002US-0385708P.
 PR PR 10-JUL-2002; 2002US-0394625P.
 PR PR 24-JUL-2002; 2002US-0398008P.
 PR PR 09-AUG-2002; 2002US-0402131P.
 PR PR 13-AUG-2002; 2002US-0402708P.
 PR PR 18-SEP-2002; 2002US-0411426P.
 PR PR 02-OCT-2002; 2002US-0414984P.
 PR PR 11-OCT-2002; 2002US-0417611P.
 PR PR 23-OCT-2002; 2002US-0420246P.
 PR PR 05-NOV-2002; 2002US-0423623P.
 PA PA (HUMA-) HUMAN GENOME SCI INC.
 PA PA (DELLZ-) DELTA BIOTECHNOLOGY LTD.
 PA PA (PRIN-) PRINCIPIA PHARM CORP.
 XX XX Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX XX WPI; 2003-598517/56.

PT PT New albumin fusion protein, useful for preparing a composition for
 PT PT treating diabetes mellitus.
 XX XX Example 4; SEQ ID NO 1523; 24pp; English.

CC CC This invention relates to a novel albumin fusion protein having albumin
 CC CC or biological activity. Human serum albumin is responsible for a
 CC CC significant proportion of the osmotic pressure of serum and also
 CC CC functions as a carrier of endogenous and exogenous ligands. The fusion of
 CC CC albumin to a therapeutic protein may increase shelf-life and stability of
 CC CC the therapeutic protein. The albumin fusion protein of the invention may
 CC CC allow production of composition with antidiabetic activity whilst the
 CC CC nucleotide sequence which encodes it may be useful for gene therapy. The
 CC CC albumin fusion protein is useful for preparing a composition for treating a
 CC CC diabetes mellitus. The present sequence is the amino acid sequence of a
 CC CC novel full-length human albumin therapeutic fusion protein of the
 CC CC invention. Note: The sequence data for this patent did not form part of
 CC CC the printed specification, but was obtained in electronic format directly
 CC CC from WIPO at ftp.wipo.int/pub/publishedpcpt sequences
 XX XX Sequence 768 AA;

QY QY 1 APPRLIDSDSIVRYLLEAKENITIGCAGHSCLSNENITPDTKNPYANKRMVEQQAA 60
 Db Db 604 APPRLIDSDSIVRYLLEAKENITIGCAGHSCLSNENITPDTKNPYANKRMVEQQAA 663
 QY QY 61 VEWQGALLSEAVLRQALYNNSOPWEPLQHVDRKAVSGRLSLTTLRALGAQEAIIS 120
 Db Db 664 VEWQGALLSEAVLRQALYNNSOPWEPLQHVDRKAVSGRLSLTTLRALGAQEAIIS 723
 QY QY 121 PPAASAPLRTTADTPKLFRVYSPFLRGKLUYGEACRTGD 165
 Db Db 724 PPDRAASAPLRTTADTPKLFRVYSPFLRGKLUYGEACRTGD 768

RESULT 135
 ID ADF16424 standard; protein; 768 AA.
 XX XX ADF16424;
 AC AC ADF16424;
 XX XX 12-FEB-2004 (first entry)
 DT DT Human albumin therapeutic fusion protein SeqID1521.
 DE DE albumin fusion protein; albumin activity; human serum albumin;
 KW KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 KW KW gene therapy; diabetes mellitus; human.
 OS OS Chimeric.
 OS OS Homo sapiens.
 XX XX WO2003060071-A2.
 PN PN 09-AUG-2002; 2002US-0402131P.
 XX XX PD 24-JUL-2003.
 PR PR 23-DEC-2002; 2002WO-US040891.
 XX XX PR 21-DEC-2001; 2001US-0341811P.
 PR PR 24-JAN-2002; 2002US-035038P.
 PR PR 28-JAN-2002; 2002US-0311360P.
 PR PR 28-FEB-2002; 2002US-0359370P.
 PR PR 02-OCT-2002; 2002US-0367500P.
 PR PR 11-OCT-2002; 2002US-0417611P.
 PR PR 09-APR-2002; 2002US-0370227P.
 PR PR 10-MAY-2002; 2002US-0378950P.
 PR PR 24-MAY-2002; 2002US-0382617P.
 PR PR 28-MAY-2002; 2002US-0383123P.
 PR PR 05-JUN-2002; 2002US-0385708P.
 PR PR 10-JUL-2002; 2002US-0394625P.
 PR PR 24-JUL-2002; 2002US-0398008P.
 PR PR 09-AUG-2002; 2002US-0402131P.
 PR PR 13-AUG-2002; 2002US-0402708P.
 PR PR 18-SEP-2002; 2002US-0411360P.
 PR PR 02-OCT-2002; 2002US-0414984P.
 PR PR 11-OCT-2002; 2002US-0417611P.
 PR PR 05-NOV-2002; 2002US-0423623P.
 PA PA (HUMA-) HUMAN GENOME SCI INC.
 PA PA (DELLZ-) DELTA BIOTECHNOLOGY LTD.
 PA PA (PRIN-) PRINCIPIA PHARM CORP.
 XX XX Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
 XX XX WPI; 2003-598517/56.

PT PT New albumin fusion protein, useful for preparing a composition for
 PT PT treating diabetes mellitus.
 XX XX Example 4; SEQ ID NO 1521; 24pp; English.

CC CC This invention relates to a novel albumin fusion protein having albumin
 or biological activity. Human serum albumin is responsible for a

CC Significant proportion of the osmotic pressure of serum and also
 CC functions as a carrier of endogenous and exogenous ligands. The fusion of
 CC albumin to a therapeutic protein may increase shelf-life and stability of
 CC the therapeutic protein. The albumin fusion protein of the invention may
 CC allow production of compositions with antidiabetic activity whilst the
 CC nucleotide sequence which encodes it may be useful for gene therapy. The
 CC albumin fusion protein is useful for preparing a composition for treating
 CC diabetes mellitus. The present sequence is the amino acid sequence of a
 CC novel full-length human albumin therapeutic fusion protein of the
 CC invention. Note: The sequence data for this patent did not form part of
 CC the printed specification, but was obtained in electronic format directly
 CC from WIPO at ftp://wipo.int/pub/publishedpat_sequences

XX Sequence 768 AA;

PR Query Match 100.0%; Score 846; DB 7; Length 768;

PR Best Local Similarity 100.0%; Pred. No. 2.1e-85; Mismatches 0; Indels 0; Gaps 0;

PR Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

PR QY 1 APPRLICDSRVLERYLLEAKERENITGCAEHCSLENITVPTDKVNFYAWKRMEVGQQA 60

PR Db 604 APPRLICDSRVLERYLLEAKERENITGCAEHCSLENITVPTDKVNFYAWKRMEVGQQA 663

PR QY 61 VEWMOGLAISLEAVIRGQALLIVNSSQWPFLQHDKAVSGLSLTTLRAIGQKEAIS 120

PR Db 664 VEWMOGLAISLEAVIRGQALLIVNSSQWPFLQHDKAVSGLSLTTLRAIGQKEAIS 723

PR QY 121 PPDAAASAPLRTTADTPRKLFRVSNPLRGKLYTGACRTGD 165

PR Db 724 PPDAAASAPLRTTADTPRKLFRVSNPLRGKLYTGACRTGD 768

RESULT 136

PR ADP16563

PR ID ADP16563 standard; protein; 768 AA.

PR AC XX

PR AC ADP16563;

PR DT 12-FEB-2004 (first entry)

PR XX Human albumin therapeutic fusion protein SeqID1660.

PR XX albumin fusion protein; albumin activity; human serum albumin;
 PR KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 PR KW gene therapy; diabetes mellitus; human.

PR OS XX Chimeric.

PR OS Homo sapiens.

PR XX WO2003060071-A2.

PR XX PD 24-JUL-2003.

PR PP 23-DEC-2002; 2002WO-US040891.

PR XX PR 21-DEC-2001; 2001US-0341811P.

PR PR 24-JAN-2002; 2002US-0350358P.

PR PR 28-JAN-2002; 2002US-0351340P.

PR PR 26-FEB-2002; 2002US-0359370P.

PR PR 28-FEB-2002; 2002US-0360000P.

PR PR 27-MAR-2002; 2002US-0367500P.

PR PR 08-APR-2002; 2002US-0370227P.

PR PR 10-MAY-2002; 2002US-0378950P.

PR PR 24-MAY-2002; 2002US-0382617P.

PR PR 28-MAY-2002; 2002US-0383123P.

PR PR 05-JUN-2002; 2002US-0385708P.

PR PR 10-JUL-2002; 2002US-0394625P.

PR PR 24-JUL-2002; 2002US-0398008P.

PR PR 09-AUG-2002; 2002US-0402131P.

PR PR 13-AUG-2002; 2002US-0402108P.

PR PR 18-SEP-2002; 2002US-0411355P.

PR PR 18-SEP-2002; 2002US-0411426P.

PR PR 02-OCT-2002; 2002US-0414984P.

PR Sequence 768 AA;

PR Query Match 100.0%; Score 846; DB 7; Length 768;

PR Best Local Similarity 100.0%; Pred. No. 2.1e-85; Mismatches 0; Indels 0; Gaps 0;

PR Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

PR QY 1 APPRLICDSRVLERYLLEAKERENITGCAEHCSLENITVPTDKVNFYAWKRMEVGQQA 60

PR Db 604 APPRLICDSRVLERYLLEAKERENITGCAEHCSLENITVPTDKVNFYAWKRMEVGQQA 663

PR QY 61 VEWMOGLAISLEAVIRGQALLIVNSSQWPFLQHDKAVSGLSLTTLRAIGQKEAIS 120

PR Db 664 VEWMOGLAISLEAVIRGQALLIVNSSQWPFLQHDKAVSGLSLTTLRAIGQKEAIS 723

PR QY 121 PPDAAASAPLRTTADTPRKLFRVSNPLRGKLYTGACRTGD 165

PR Db 724 PPDAAASAPLRTTADTPRKLFRVSNPLRGKLYTGACRTGD 768

PR RESULT 137

PR ADP15091

PR ID ADP15091 standard; protein; 769 AA.

PR AC XX

PR AC ADP15091;

PR DT 12-FEB-2004 (first entry)

PR XX Human albumin therapeutic fusion protein SeqID387.

PR XX albumin fusion protein; albumin activity; human serum albumin;
 PR KW serum osmotic pressure; shelf-life; stability; antidiabetic;
 PR KW gene therapy; diabetes mellitus; human.

PR OS XX Chimeric.

PR OS Homo sapiens.

PR XX WO2003060071-A2.

PR XX PD 24-JUL-2003.

PR PP 23-DEC-2002; 2002WO-US040891.

XX
PR 21-DEC-2001; 2001US-0341811P.
PR 24-JAN-2002; 2002US-0350358P.
PR 28-JAN-2002; 2002US-0351360P.
PR 26-FEB-2002; 2002US-0359370P.
PR 27-MAR-2002; 2002US-0367500P.
PR 08-APR-2002; 2002US-0370227P.
PR 10-MAY-2002; 2002US-0378950P.
PR 24-MAY-2002; 2002US-0382617P.
PR 28-MAY-2002; 2002US-0383123P.
PR 05-JUN-2002; 2002US-0385703P.
PR 10-JUL-2002; 2002US-0394625P.
PR 24-JUL-2002; 2002US-0398008P.
PR 09-AUG-2002; 2002US-0402131P.
PR 13-AUG-2002; 2002US-0402708P.
PR 18-SEP-2002; 2002US-0411355P.
PR 18-SEP-2002; 2002US-0411426P.
PR 02-OCT-2002; 2002US-0414984P.
PR 11-OCT-2002; 2002US-0417611P.
PR 23-OCT-2002; 2002US-0420246P.
PR 05-NOV-2002; 2002US-0423623P.
PA (HUMA-) HUMAN GENOME SCI INC.
PA (DELBZ) DELTA BIOTECHNOLOGY LTD.
PA (PRIN-) PRINCIPIA PHARM CORP.
XX
PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
DR WPI; 2003-598517/56.
XX
PT New albumin fusion protein, useful for preparing a composition for
PT treating diabetes mellitus.
XX
Example 4; SEQ ID NO 387; 24pp; English.
XX
CC This invention relates to a novel albumin fusion protein having albumin
CC or biological activity. Human serum albumin is responsible for a
CC significant proportion of the osmotic pressure of serum and also
functions as a carrier of endogenous and exogenous ligands. The fusion of
albumin to a therapeutic protein may increase shelf-life and stability of
the therapeutic protein. The albumin fusion protein of the invention may
allow production of compositions with antidiabetic activity whilst the
nucleotide sequence which encodes it may be useful for gene therapy. The
albumin fusion protein is useful for preparing a composition for treating
diabetes mellitus. The present sequence is the amino acid sequence of a
novel full-length human albumin therapeutic fusion protein of the
invention. Note: The sequence data for this patent did not form part of
the printed specification, but was obtained in electronic format directly
from WIPO at ftp://wipo.int/pub/publishedpat_sequences
SQ Sequence 769 AA;

Query Match 100.0%; Score 846; DB 7; Length 769;
Best Local Similarity 100.0%; Pred. No. 2.1e-85; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKNAENITTCGAEHCGLNENTTVPDKTKNFYAWKRMVEVGQA 60
DB 20 APPRLICDSRVRLERYLLEAKNAENITTCGAEHCGLNENTTVPDKTKNFYAWKRMVEVGQA 79
QY 61 VEVNOGLALIISEAVERGQALVNNSOPWELPLQHVDAVKAVSGLSLTTLRAGKQEAIS 120
DB 80 VEVNOGLALIISEAVERGQALVNNSOPWELPLQHVDAVKAVSGLSLTTLRAGKQEAIS 139
QY 121 PPDASAPRITIATDPRKFLPRYSNFGKGKLYTGACRGD 165
DB 140 PPDASAPRITIATDPRKFLPRYSNFGKGKLYTGACRGD 184
XX
RESULT 138
ID ADF15082
ID ADF15082 standard; protein; 777 AA.

XX
PR XX
AC ADP15082;
XX DT 12-FEB-2004 (first entry)
XX DE Human albumin therapeutic fusion protein SeqID378.
XX KW albumin fusion protein; albumin activity; human serum albumin;
serum osmotic pressure; shelf-life; stability; antidiabetic;
gene therapy; diabetes mellitus; human.
XX OS Homo sapiens.
XX PN WO2003060071-A2.
XX PD 24-JUL-2003.
XX PR XX
PR 23-DEC-2002; 2002WO-US040891.
PR 21-DEC-2001; 2001US-0341811P.
PR 24-JAN-2002; 2002US-0350358P.
PR 28-JAN-2002; 2002US-0351360P.
PR 26-FEB-2002; 2002US-0359370P.
PR 28-FEB-2002; 2002US-0360000P.
PR 27-MAR-2002; 2002US-0367500P.
PR 08-APR-2002; 2002US-0370227P.
PR 10-MAY-2002; 2002US-0378950P.
PR 24-MAY-2002; 2002US-0402131P.
PR 28-MAY-2002; 2002US-0383123P.
PR 05-JUN-2002; 2002US-0385703P.
PR 10-JUL-2002; 2002US-0394625P.
PR 09-AUG-2002; 2002US-0402131P.
PR 13-AUG-2002; 2002US-0402708P.
PR 18-SEP-2002; 2002US-0411355P.
PR 18-SEP-2002; 2002US-0411426P.
PR 02-OCT-2002; 2002US-0414984P.
PR 11-OCT-2002; 2002US-0417611P.
PR 23-OCT-2002; 2002US-0420246P.
PR 05-NOV-2002; 2002US-0423623P.
PA (HUMA-) HUMAN GENOME SCI INC.
PA (DELBZ) DELTA BIOTECHNOLOGY LTD.
PA (PRIN-) PRINCIPIA PHARM CORP.
XX
PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;
DR WPI; 2003-598517/56.
XX
PT New albumin fusion protein, useful for preparing a composition for
PT treating diabetes mellitus.
XX
Example 4; SEQ ID NO 378; 24pp; English.
XX
CC This invention relates to a novel albumin fusion protein having albumin
CC or biological activity. Human serum albumin is responsible for a
CC significant proportion of the osmotic pressure of serum and also
functions as a carrier of endogenous and exogenous ligands. The fusion of
albumin to a therapeutic protein may increase shelf-life and stability of
the therapeutic protein. The albumin fusion protein of the invention may
allow production of compositions with antidiabetic activity whilst the
nucleotide sequence which encodes it may be useful for gene therapy. The
albumin fusion protein is useful for preparing a composition for treating
diabetes mellitus. The present sequence is the amino acid sequence of a
novel full-length human albumin therapeutic fusion protein of the
invention. Note: The sequence data for this patent did not form part of
the printed specification, but was obtained in electronic format directly
from WIPO at ftp://wipo.int/pub/publishedpat_sequences
SQ Sequence 777 AA;

Query Match 100.0%; Score 846; DB 7; Length 777;

Best Local Similarity 100.0%; **Pred.** No. 2.1e-85; **Matches** 165; **Conservative** 0; **Mismatches** 0; **Indels** 0; **Gaps** 0; **ID** ADF15078; **XX** ADP15078; **AC** ADP15078; **DT** 12-FEB-2004 (first entry) **DE** Human albumin therapeutic fusion protein SeqID374. **KW** albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human. **XX** Chimeric. **OS** Homo sapiens. **XX** WO2003060071-A2. **XX** PD 24-JUL-2003. **XX** PF 23-DEC-2002; 2002WO-US040891. **PR** 21-DEC-2001; 2001US-0341811P. **PR** 24-JAN-2002; 2002US-0350388P. **PR** 28-JAN-2002; 2002US-0351360P. **PR** 26-FEB-2002; 2002US-0359370P. **PR** 28-FEB-2002; 2002US-0360000P. **PR** 08-APR-2002; 2002US-0367500P. **PR** 10-MAY-2002; 2002US-0378950P. **PR** 24-MAY-2002; 2002US-0382617P. **PR** 28-MAY-2002; 2002US-0383123P. **PR** 05-JUN-2002; 2002US-0385708P. **PR** 10-JUL-2002; 2002US-0394625P. **PR** 24-JUL-2002; 2002US-039908P. **PR** 09-AUG-2002; 2002US-0402131P. **PR** 13-AUG-2002; 2002US-0402708P. **PR** 18-SEP-2002; 2002US-041126P. **PR** 02-OCT-2002; 2002US-0414984P. **PR** 11-OCT-2002; 2002US-0417611P. **PR** 23-OCT-2002; 2002US-0420246P. **PR** 05-NOV-2002; 2002US-0423623P. **XX** (HUMA-) HUMAN GENOME SCI INC. **PA** (DELB-) DELTA BIOTECHNOLOGY LTD. **PA** (PRIN-) PRINCIPIA PHARM CORP. **XX** Ballance DJ, Turner AJ, Rosen CA, Haseltine WA; **XX** WPI; 2003-59851/756.

Best Local Similarity 100.0%; **Pred.** No. 2.1e-85; **Matches** 165; **Conservative** 0; **Mismatches** 0; **Indels** 0; **Gaps** 0; **ID** QY 1 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 60 **Db** 28 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 87 **QY** 61 VEWMOGLALLSSEAVLRGQALLVNSQWEPLQLHVDAVSGRSLSLTLLRALGAOKEAI 120 **Db** 88 VEWMOGLALLSSEAVLRGQALLVNSQWEPLQLHVDAVSGRSLSLTLLRALGAOKEAI 147 **QY** 121 PPDASASAPERTITADTPRKLFPRVYSNPLRGKLYTGEACRTGD 165 **Db** 148 PPDASASAPERTITADTPRKLFPRVYSNPLRGKLYTGEACRTGD 192

RESULT 139 **SQ** Sequence 777 AA; **ID** ADF15078 **XX** ADP15078 standard; protein: 777 AA. **Query Match** Best Local Similarity 100.0%; Score 846; DB 7; Length 777; **Matches** 165; **Conservative** 0; **Mismatches** 0; **Indels** 0; **Gaps** 0; **ID** QY 1 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 60 **Db** 28 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 87 **QY** 61 VEWMOGLALLSSEAVLRGQALLVNSQWEPLQLHVDAVSGRSLSLTLLRALGAOKEAI 120 **Db** 88 VEWMOGLALLSSEAVLRGQALLVNSQWEPLQLHVDAVSGRSLSLTLLRALGAOKEAI 147 **QY** 121 PPDASASAPERTITADTPRKLFPRVYSNPLRGKLYTGEACRTGD 165 **Db** 148 PPDASASAPERTITADTPRKLFPRVYSNPLRGKLYTGEACRTGD 192

RESULT 140 **SQ** Sequence 777 AA; **ID** ADF15075 **XX** ADP15075 standard; protein: 777 AA. **Query Match** Best Local Similarity 100.0%; Score 846; DB 7; Length 777; **Matches** 165; **Conservative** 0; **Mismatches** 0; **Indels** 0; **Gaps** 0; **ID** QY 1 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 60 **Db** 28 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 87 **DE** Human albumin therapeutic fusion Protein SeqID371. **KW** albumin fusion protein; albumin activity; human serum albumin; serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human. **XX** Chimeric. **OS** Homo sapiens. **XX** WO2003060071-A2. **XX** PD 24-JUL-2003. **XX** PF 23-DEC-2002; 2002WO-US040891. **PR** 21-DEC-2001; 2001US-0341811P. **PR** 24-JAN-2002; 2002US-0350388P. **PR** 28-JAN-2002; 2002US-0351360P. **PR** 26-FEB-2002; 2002US-0359370P. **PR** 28-FEB-2002; 2002US-0360000P. **PR** 08-APR-2002; 2002US-0367500P. **PR** 10-MAY-2002; 2002US-0378950P. **PR** 24-MAY-2002; 2002US-0382617P. **PR** 28-MAY-2002; 2002US-0383123P. **PR** 05-JUN-2002; 2002US-0385708P. **PR** 10-JUL-2002; 2002US-0394625P. **PR** 24-JUL-2002; 2002US-039908P. **PR** 09-AUG-2002; 2002US-0402131P. **PR** 13-AUG-2002; 2002US-0402708P. **PR** 18-SEP-2002; 2002US-041126P. **PR** 02-OCT-2002; 2002US-0414984P. **PR** 11-OCT-2002; 2002US-0417611P. **PR** 23-OCT-2002; 2002US-0420246P. **PR** 05-NOV-2002; 2002US-0423623P. **XX** (HUMA-) HUMAN GENOME SCI INC. **PA** (DELB-) DELTA BIOTECHNOLOGY LTD. **PA** (PRIN-) PRINCIPIA PHARM CORP. **XX** Ballance DJ, Turner AJ, Rosen CA, Haseltine WA; **XX** WPI; 2003-59851/756.

Best Local Similarity 100.0%; **Pred.** No. 2.1e-85; **Matches** 165; **Conservative** 0; **Mismatches** 0; **Indels** 0; **Gaps** 0; **ID** QY 1 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 60 **Db** 28 APPRLICDSVRLERYLLEAKERENITTGCAEBCSLENNTIVPDTKVNFYAWKRMVEGQQA 87 **CC** This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is the amino acid sequence of a novel full-length human albumin therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/publishedpat_sequences

SQ	Sequence 777 AA;	Query Match	100.0%	Score 846;	DB 7;	Length 777;
ID	ADP15079 standard; protein; 777 AA.	Best Local Similarity	100.0%	Pred. No.	2.1e-85;	
AC	ADP15079;	Matches	165;	Conservative	0;	Mismatches
XX	DT 12-FEB-2004 (first entry)	Indels	0;	Gaps	0;	
XX	DE Human albumin therapeutic fusion protein SeqID375.					
XX	KW albumin fusion protein; albumin activity; human serum albumin; gene therapy; shelf-life; stability; antidiabetic; human.					
KW	serum osmotic pressure; shelf-life; stability; antidiabetic; gene therapy; diabetes mellitus; human.					
OS	Chimeric.					
OS	Homo sapiens.					
XX	PN WO2003060071-A2.					
XX	PD 24-JUL-2003.					
XX	PP 23-DEC-2002; 2002WO-US040891.					
XX	PR 21-DEC-2001; 2001US-0341811P.					
PR	24-JAN-2002; 2002US-0350358P.					
PR	28-JAN-2002; 2002US-0351360P.					
PR	28-FEB-2002; 2002US-0359370P.					
PR	27-MAR-2002; 2002US-0367500P.					
PR	08-APR-2002; 2002US-0370221P.					
PR	10-MAY-2002; 2002US-0378950P.					
PR	24-MAY-2002; 2002US-0382617P.					
PR	28-MAY-2002; 2002US-0383123P.					
PR	05-JUN-2002; 2002US-0385700P.					
PR	10-JUL-2002; 2002US-0394625P.					
PR	24-JUL-2002; 2002US-0398008P.					
PR	09-AUG-2002; 2002US-0402131P.					
PR	13-AUG-2002; 2002US-0402700P.					
PR	18-SEP-2002; 2002US-0411355P.					
PR	02-OCT-2002; 2002US-0414984P.					
PR	11-OCT-2002; 2002US-0417611P.					
PR	23-OCT-2002; 2002US-0420246P.					
PR	05-NOV-2002; 2002US-0423623P.					
XX	(HUMA-) HUMAN GENOME SCI INC.					
PA	(DELL-) DELTA BIOTECHNOLOGY LTD.					
PA	(PRIN-) PRINCIPIA PHARM CORP.					
PR	XX WO2003060071-A2.					
XX	PD 24-JUL-2003.					
XX	PP 23-DEC-2002; 2002WO-US040891.					
XX	PR 21-DEC-2001; 2001US-0341811P.					
PR	24-JAN-2002; 2002US-0350358P.					
PR	28-JAN-2002; 2002US-0351360P.					
PR	26-FEB-2002; 2002US-0359370P.					
PR	28-FEB-2002; 2002US-0360000P.					
PR	27-MAR-2002; 2002US-0367500P.					
PR	08-APR-2002; 2002US-0370221P.					
PR	10-MAY-2002; 2002US-0378950P.					
PR	24-MAY-2002; 2002US-0382617P.					
PR	28-MAY-2002; 2002US-0383123P.					
PR	05-JUN-2002; 2002US-0385700P.					
PR	10-JUL-2002; 2002US-0394625P.					
PR	24-JUL-2002; 2002US-0398008P.					
PR	09-AUG-2002; 2002US-0402131P.					
PR	13-AUG-2002; 2002US-0402700P.					
PR	18-SEP-2002; 2002US-0411355P.					
PR	05-NOV-2002; 2002US-0423623P.					
XX	(HUMA-) HUMAN GENOME SCI INC.					
PA	(DELL-) DELTA BIOTECHNOLOGY LTD.					
PA	(PRIN-) PRINCIPIA PHARM CORP.					
PT	XX WO2003060071-A2.					
PT	Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;					
PT	XX WPI; 2003-598517/56.					
PT	PT New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.					
PS	Example 4; SEQ ID NO 375; 24pp; English.					
XX	This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present invention sequence is the amino acid sequence of a novel full-length human therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp.wipo.int/pub/publishedpatent_sequences					
XX	PT New albumin fusion protein, useful for preparing a composition for					

PT treating diabetes mellitus.

XX
PS Example 4; SEQ ID NO 377; 24pp; English.

This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is the amino acid sequence of a novel full-length human albumin therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp://wipo.int/pub/publishedpat_sequences

XX SQ Sequence 777 AA;

Query Match 100.0%; Score 846; DB 7; Length 777;
Best Local Similarity 100.0%; Pred. No. 2. 1e-85; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 1
1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNLYAWKRMGVQQA 60
28 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNLYAWKRMGVQQA 87
QY 61 VEWQGLALISEAVLRGQALLVNSQPWPLQLAVDKAVGSLRSLTTLRAGQKEAIS 120
Db 88 VEWQGLALISEAVLRGQALLVNSQPWPLQLAVDKAVGSLRSLTTLRAGQKEAIS 147
QY 121 PPDAASAPLRTTADTPRKLFVSYNPLRGKLYGEACRGD 165
Db 148 PPDAASAPLRTTADTPRKLFVSYNPLRGKLYGEACRGD 192

RESULT 144

ID ADF15113
ID ADF15113 standard; protein; 951 AA.

XX AC ADF15113;
XX DT 12-FEB-2004 (first entry)

DB Human albumin therapeutic fusion protein SeqID409.
XX KW albumin fusion protein; albumin activity; human serum albumin;
KW serum osmotic pressure; shelf-life; stability; antidiabetic;
KW gene therapy; diabetes mellitus; human.

XX OS Chimeric.
OS Homo sapiens.
XX PN WO2003060071-A2.

XX PD 24-JUL-2003.

PP 23 - DBC-2002; 2002WO-US040891.

XX PR 21-DEC-2001; 2001US-0341811P.

PR 24-JAN-2002; 2002US-030358P.

PR 28-JAN-2002; 2002US-0311360P.

PR 26-FEB-2002; 2002US-0359370P.

PR 28-FEB-2002; 2002US-036000P.

PR 27-MAR-2002; 2002US-0367500P.

PR 08-APR-2002; 2002US-0370227P.

PR 10-MAY-2002; 2002US-0378950P.

PR 24-MAY-2002; 2002US-0302617P.

PR 05-JUN-2002; 2002US-0333123P.

PR 10-JUL-2002; 2002US-0394625P.

PR 24-JUL-2002; 2002US-0398008P.

PR 09-AUG-2002; 2002US-0402131P.

PR 13-AUG-2002; 2002US-0402708P.

PR 18-SEP-2002; 2002US-041135P.

PR 03-OCT-2002; 2002US-0411426P.

PR 11-OCT-2002; 2002US-041761P.

PR 22-OCT-2002; 2002US-0420246P.

PR 05-NOV-2002; 2002US-0423623P.

XX PA (HUMA-) HUMAN GENOME SCI INC.

PA (DELZ-) DELTA BIOTECHNOLOGY LTD.

(PRIN-) PRINCIPIA PHARM CORP.

PI Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;

DR WPI; 2003-598517/56.

XX PT New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.

XX PS Example 4; SEQ ID NO 409; 24pp; English.

XX This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is the amino acid sequence of a novel full-length human albumin therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp://wipo.int/pub/publishedpat_sequences

XX SQ Sequence 951 AA;

Query Match 100.0%; Score 846; DB 7; Length 951;
Best Local Similarity 100.0%; Pred. No. 2. 8e-85; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 1
1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNLYAWKRMGVQQA 60
28 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNLYAWKRMGVQQA 87
QY 61 VEWQGLALIS EA VL RG Q ALL V NS Q P W PL Q L AV D KA VG S L R S LT T L R A G Q K E A I S 120
Db 88 VEWQGLALIS EA VL RG Q ALL V NS Q P W PL Q L AV D KA VG S L R S LT T L R A G Q K E A I S 147
QY 121 PPDAASAPLRTTADTPRKLFVSYNPLRGKLYGEACRGD 165
Db 148 PPDAASAPLRTTADTPRKLFVSYNPLRGKLYGEACRGD 192

RESULT 145

ID ADF15108
ID ADF15108 standard; protein; 951 AA.

XX AC ADF15108;
XX DT 12-FEB-2004 (first entry)

DB Human albumin therapeutic fusion protein SeqID404.
XX KW albumin fusion protein; albumin activity; human serum albumin;
KW serum osmotic pressure; shelf-life; stability; antidiabetic;
KW gene therapy; diabetes mellitus; human.

XX OS Chimeric.

OS Homo sapiens.

XX	PN	WO2003060071-A2.	Db	148	FPDAAASAPRPTITADTPRKFLFVSNFLRGKLYTGACRTGQ	192
PD	XX	24-JUL-2003.				
PR	XX	23-DEC-2002; 2002WO-US040891.	ID	ADFI5105	RESULT 146	
PR	XX		ID	ADFI5105 standard; protein; 954 AA.	ADFI5105;	
PR	XX		ID	ADFI5105;	ADFI5105;	
PR	XX	21-DEC-2001; 2001US-0341811P.	ID	ADFI5105	ADFI5105;	
PR	XX	20-JAN-2002; 2002US-0351360P.	ID	ADFI5105	ADFI5105;	
PR	XX	26-FEB-2002; 2002US-0359370P.	ID	ADFI5105	ADFI5105;	
PR	XX	26-FEB-2002; 2002US-0360000P.	ID	ADFI5105	ADFI5105;	
PR	XX	08-APR-2002; 2002US-0370227P.	ID	ADFI5105	ADFI5105;	
PR	XX	10-MAY-2002; 2002US-0378950P.	ID	ADFI5105	ADFI5105;	
PR	XX	24-MAY-2002; 2002US-0392617P.	ID	ADFI5105	ADFI5105;	
PR	XX	28-MAY-2002; 2002US-0393123P.	ID	ADFI5105	ADFI5105;	
PR	XX	05-JUN-2002; 2002US-0395708P.	ID	ADFI5105	ADFI5105;	
PR	XX	10-JUL-2002; 2002US-0394625P.	ID	ADFI5105	ADFI5105;	
PR	XX	09-AUG-2002; 2002US-0402131P.	ID	ADFI5105	ADFI5105;	
PR	XX	13-AUG-2002; 2002US-0402708P.	ID	ADFI5105	ADFI5105;	
PR	XX	18-SEP-2002; 2002US-0411355P.	ID	ADFI5105	ADFI5105;	
PR	XX	18-SEP-2002; 2002US-0411426P.	ID	ADFI5105	ADFI5105;	
PR	XX	05-OCT-2002; 2002US-0414984P.	ID	ADFI5105	ADFI5105;	
PR	XX	11-OCT-2002; 2002US-0417611P.	ID	ADFI5105	ADFI5105;	
PR	XX	23-OCT-2002; 2002US-0420246P.	ID	ADFI5105	ADFI5105;	
PR	XX	05-NOV-2002; 2002US-0423623P.	ID	ADFI5105	ADFI5105;	
PA	(HUMA-) HUMAN GENOME SCI INC.	(DELZ) DELTA BIOTECHNOLOGY LTD.	AC	ADFI5105;	ADFI5105;	
PA	(PRIN-) PRINCIPIA PHARM CORP.	(PRIN-) PRINCIPIA PHARM CORP.	AC	ADFI5105;	ADFI5105;	
PT	New albumin fusion protein, useful for preparing a composition for treating diabetes mellitus.		AC	ADFI5105;	ADFI5105;	
PT	Ballance DJ, Turner AJ, Rosen CA, Haseltine WA;		AC	ADFI5105;	ADFI5105;	
PI	WPI; 2003-598517/56.		AC	ADFI5105;	ADFI5105;	
XX	This invention relates to a novel albumin fusion protein having albumin or biological activity. Human serum albumin is responsible for a significant proportion of the osmotic pressure of serum and also functions as a carrier of endogenous and exogenous ligands. The fusion of albumin to a therapeutic protein may increase shelf-life and stability of the therapeutic protein. The albumin fusion protein of the invention may allow production of compositions with antidiabetic activity whilst the nucleotide sequence which encodes it may be useful for gene therapy. The albumin fusion protein is useful for preparing a composition for treating diabetes mellitus. The present sequence is the amino acid sequence of a novel full-length human albumin therapeutic fusion protein of the invention. Note: The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp://ftp.wipo.int/pub/publishedpat_sequences		AC	ADFI5105;	ADFI5105;	
SQ	Sequence 951 AA;		AC	ADFI5105;	ADFI5105;	
Query	Match	100.0%; Score 846; DB 7; Length 951;	AC	ADFI5105;	ADFI5105;	
Best	Local Similarity	100.0%; Pred. No. 2.8e-85;	AC	ADFI5105;	ADFI5105;	
Matches	165; Conservative	0; Mismatches 0; Indels 0; Gaps 0;	AC	ADFI5105;	ADFI5105;	
QY	1 APPRLICDSRVLERVYLEAKAENITTGCAEHCSLENITVPDTKVNFKWAKMVGQQA 60	2 APPRLICDSRVLERVYLEAKAENITTGCAEHCSLENITVPDTKVNFKWAKMVGQQA 87	CC	ADFI5105;	ADFI5105;	
Db	61 VEVNGGLALLSEAVVRGQQLVNNSOPWEPQLQHDKAVSGLSRSTLRLAGKEAT 120	88 VEVNGGLALLSEAVVRGQQLVNNSOPWEPQLQHDKAVSGLSRSTLRLAGKEAT 147	CC	ADFI5105;	ADFI5105;	
Oy	121 PPDAAASAPRPTITADTPRKFLFVSNFLRGKLYTGACRTGQ 165		CC	ADFI5105;	ADFI5105;	

CC the printed specification, but was obtained in electronic format directly
CC from WIPO at ftp://ftp.wipo.int/pub/publishedpt_sequences

XX
SQ Sequence 954 AA;

Query Match 100.0%; Score 846; DB 7; Length 954;
Best Local Similarity 100.0%; Pred. No. 2. 9e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRICOSRVERLYLRAKEAENITGCAEHCSLENENTPDKYKVNFTAKRMEVGQA 60
Db 790 APPRICOSRVERLYLRAKEAENITGCAEHCSLENENTPDKYKVNFTAKRMEVGQA 849

Qy 61 VEWQGLAISSEAVLRGQALVNSQPMELQLQHVDKAVSGRLSITLRLGAQKAIS 120
Db 850 VEWQGLAISSEAVLRGQALVNSQPMELQLQHVDKAVSGRLSITLRLGAQKAIS 909

Qy 121 PPDASAAPLRTTADTFRKLFRTYSNPRGKULYTGEACRTGD 165
Db 910 PPDASAAPLRTTADTFRKLFRTYSNPRGKULYTGEACRTGD 954

Search completed: March 1, 2006, 10:23:29
Job time : 196 secs

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 GenCore version 5.1.7

OM protein - protein search, using sw model
 Run on: March 1, 2006, 10:20:21 ; Search time 65 Seconds
 (without alignments)
 1060.644 Million cell updates/sec

Title: US-10-706-701-1
 Perfect score: 846
 Sequence: 1 APPRLICDSVRLERYLRLBAK. SNPLRGKLUKYTGACRGD 165

Scoring table: BLOSUM62
 Gapop 10.0 , Gapext 0.5
 Searched: 1867569 seqs, 417829326 residues

Total number of hits satisfying chosen parameters: 102
 Minimum DB seq length: 0
 Maximum DB seq length: 200000000

Post-processing: Minimum Match 100%
 Maximum Match 100%
 Listing first 500 summaries

Database : Published Applications AA_Main:
 1: /cgn2_6/ptodata/1/pupaa/US07_PUBCOMB.pep:
 2: /cgn2_6/ptodata/1/pupaa/US08_PUBCOMB.pep:
 3: /cgn2_6/ptodata/1/pupaa/US09_PUBCOMB.pep:
 4: /cgn2_6/ptodata/1/pupaa/US10A_PUBCOMB.pep:
 5: /cgn2_6/ptodata/1/pupaa/US10B_PUBCOMB.pep:
 6: /cgn2_6/ptodata/1/pupaa/US11_PUBCOMB.pep:

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match Length	DB ID	Description
1	846	100.0	165	3 US-09-853-731-1 Sequence 1, Appli
2	846	100.0	165	3 US-09-945-517-1 Sequence 1, Appli
3	846	100.0	165	4 US-10-014-363-1 Sequence 1, Appli
4	846	100.0	165	4 US-10-241-356-1 Sequence 1, Appli
5	846	100.0	165	4 US-10-293-551-1 Sequence 1, Appli
6	846	100.0	165	4 US-10-411-037-73 Sequence 1, Appli
7	846	100.0	165	4 US-10-411-026-73 Sequence 1, Appli
8	846	100.0	165	4 US-10-410-962-73 Sequence 1, Appli
9	846	100.0	165	4 US-10-411-049-73 Sequence 1, Appli
10	846	100.0	165	4 US-10-634-477-1 Sequence 1, Appli
11	846	100.0	165	4 US-10-410-930-73 Sequence 1, Appli
12	846	100.0	165	4 US-10-410-997-73 Sequence 1, Appli
13	846	100.0	165	4 US-10-411-012-73 Sequence 1, Appli
14	846	100.0	165	4 US-10-410-913-2 Sequence 1, Appli
15	846	100.0	165	4 US-10-780-297-1 Sequence 1, Appli
16	846	100.0	165	4 US-10-706-701-1 Sequence 1, Appli
17	846	100.0	165	5 US-10-410-90-73 Sequence 1, Appli
18	846	100.0	165	5 US-10-410-889-73 Sequence 1, Appli
19	846	100.0	165	6 US-11-013-560-1 Sequence 1, Appli
20	846	100.0	166	3 US-09-853-731-2 Sequence 1, Appli
21	846	100.0	166	4 US-10-014-363-2 Sequence 1, Appli
22	846	100.0	166	4 US-10-241-356-2 Sequence 1, Appli
23	846	100.0	166	4 US-10-93-551-2 Sequence 1, Appli
24	846	100.0	166	4 US-10-400-377-2 Sequence 1, Appli
25	846	100.0	166	4 US-10-400-708-2 Sequence 1, Appli
26	846	100.0	166	4 US-10-298-148-2 Sequence 1, Appli
27	846	100.0	166	4 US-10-360-101-227 Sequence 1, Appli

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Sequence 367, Ap
Sequence 371, Ap
Sequence 374, Ap
Sequence 375, Ap
Sequence 377, Ap
Sequence 378, Ap
Sequence 404, Ap

RESULT 1
US-09-853-731-1
; Sequence 1, Application US/09853731
; Patent No. US20020031841A1
; GENERAL INFORMATION:
; APPLICANT: Papadimitriou, Apollon
; TITLE OF INVENTION: Erythropoietin Composition
; FILE REFERENCE: 20619 US
; CURRENT APPLICATION NUMBER: US/09/853, 731
; CURRENT FILING DATE: 2001-05-11
; PRIORITY APPLICATION NUMBER: EP/00110355.5
; PRIORITY FILING DATE: 2000-05-15
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.0
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-853-731-1

Query Match 100.0%; Score 846; DB 3; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRPLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
Db 1 APPRLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
QY 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120
Db 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120
QY 121 PPDASAAPRITTADETRKLFRVSNFLRGKLYTGEACRTGD 165
Db 121 PPDASAAPRITTADETRKLFRVSNFLRGKLYTGEACRTGD 165

RESULT 2
US-09-945-517-1
; Sequence 1, Application US/09945517
; Publication No. US2003010496A1
; GENERAL INFORMATION:
; APPLICANT: Li, Tiansheng
; APPLICANT: Chang, Byeong
; APPLICANT: Sloey, Christopher
; TITLE OF INVENTION: L-METHIONINE AS A STABILIZER FOR NSP/EPO IN HSA-FREE FORMULATION
; FILE REFERENCE: A-803
; CURRENT APPLICATION NUMBER: US/09/945, 517
; CURRENT FILING DATE: 2001-08-30
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.0
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-945-517-1

Query Match 100.0%; Score 846; DB 3; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRPLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
Db 1 APPRLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
QY 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120

RESULT 3
US-10-014-363-1
; Sequence 1, Application US/10014363
; Publication No. US20020115833A1
; GENERAL INFORMATION:
; APPLICANT: Burg, Josef
; APPLICANT: Engel, Alfred
; APPLICANT: Franze, Reinhard
; APPLICANT: Hilger, Bernd
; APPLICANT: Schurig, Hartmut Ernst
; APPLICANT: Tischer, Wilhelm
; APPLICANT: Wozny, Manfred
; TITLE OF INVENTION: Erythropoietin Conjugates
; FILE REFERENCE: Case 2005
; CURRENT APPLICATION NUMBER: US/10/014, 363
; CURRENT FILING DATE: 2001-12-11
; NUMBER OF SEQ ID NOS: 5
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-014-363-1

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRPLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
Db 1 APPPLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
QY 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120
Db 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120
QY 121 PPDASAAPRITTADETRKLFRVSNFLRGKLYTGEACRTGD 165
Db 121 PPDASAAPRITTADETRKLFRVSNFLRGKLYTGEACRTGD 165

RESULT 4
US-10-241-356-1
; Sequence 1, Application US/10241356
; Publication No. US20030077753A1
; GENERAL INFORMATION:
; APPLICANT: TISCHER, WILHELM
; FILE REFERENCE: 20971
; CURRENT APPLICATION NUMBER: US/10/241, 356
; CURRENT FILING DATE: 2002-09-11
; PRIORITY APPLICATION NUMBER: EP 01122555.4
; PRIORITY FILING DATE: 2001-09-25
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-241-356-1

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APRPLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
Db 1 APPRLCDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNFKWAKMVEGQOA 60
QY 61 VEWMOGLALISEAVLRQQLVNSQWPEDLQLHVDKAVSGLRSLSLTLLRAQKEAIS 120

QY 1 APPLICDSRVLERYLLEAKRENTITGCAEHCSLENITVDPDKVNFYAWKRMENGQA 60

QY 61 VEWQGLALLSEAVRGQALVNNSQPWPQLHQDKAVSGRLSLTLLRAIGQKAIS 120

Db 61 VEWQGLALLSEAVRGQALVNNSQPWPQLHQDKAVSGRLSLTLLRAIGQKAIS 120

Db 121 PPDASAPRLTTADTFRKLFPRVYSNPLRGKLUYTGEACRTGD 165

RESULT 5
US-10-293-551-1
; Sequence 1, Application US/10293551
; Publication No. US20030120045A1
; GENERAL INFORMATION:
; APPLICANT: Ballon, Pascal
; TITLE OF INVENTION: ERYTHROPOIETIN CONJUGATES
; FILE NUMBER: 1097 nonprovisional
; CURRENT APPLICATION NUMBER: US/10/293,551
; CURRENT FILING DATE: 2002-11-14
; PRIOR APPLICATION NUMBER: US/09/604,938
; PRIOR FILING DATE: 2000-06-27
; PRIOR APPLICATION NUMBER: 60/166,151
; PRIOR FILING DATE: 1999-11-17
; PRIOR APPLICATION NUMBER: 60/151,548
; PRIOR FILING DATE: 1999-08-13
; PRIOR APPLICATION NUMBER: 60/150,225
; PRIOR FILING DATE: 1999-08-23
; PRIOR APPLICATION NUMBER: 60/142,254
; PRIOR FILING DATE: 1999-07-02
; NUMBER OF SEQ ID NOS: 3
; SOFTWARE: Patentin Ver. 2.1
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-293-551-1

Query Match Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPLICDSRVLERYLLEAKRENTITGCAEHCSLENITVDPDKVNFYAWKRMENGQA 60

Db 1 APPLICDSRVLERYLLEAKRENTITGCAEHCSLENITVDPDKVNFYAWKRMENGQA 60

QY 61 VEWQGLALLSEAVRGQALVNNSQPWPQLHQDKAVSGRLSLTLLRAIGQKAIS 120

Db 61 VEWQGLALLSEAVRGQALVNNSQPWPQLHQDKAVSGRLSLTLLRAIGQKAIS 120

QY 121 PPDASAPRLTTADTFRKLFPRVYSNPLRGKLUYTGEACRTGD 165

Db 121 PPDASAPRLTTADTFRKLFPRVYSNPLRGKLUYTGEACRTGD 165

RESULT 6
US-10-411-037-73
; Sequence 73, Application US/10411037
; Publication No. US2004006391A1
; GENERAL INFORMATION:
; APPLICANT: Neese Technologies, Inc.
; APPLICANT: Derees, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; TITLE OF INVENTION: PROTEIN REMODELING METHODS AND PROTEINS/PEPTIDES PRODUCED BY THE
; FILE REFERENCE: 040853-01-5053
; CURRENT APPLICATION NUMBER: US/10/411,026
; PRIOR FILING DATE: 2003-04-09
; PRIOR APPLICATION NUMBER: US 60/328,523
; PRIOR FILING DATE: 2001-10-0
; PRIOR APPLICATION NUMBER: US 60/344,692
; PRIOR FILING DATE: 2001-10-19
; PRIOR APPLICATION NUMBER: US 60/387,292
; PRIOR FILING DATE: 2002-06-07
; PRIOR APPLICATION NUMBER: US 60/391,777
; PRIOR FILING DATE: 2002-06-25
; PRIOR APPLICATION NUMBER: US 60/396,594
; PRIOR FILING DATE: 2002-07-17
; PRIOR APPLICATION NUMBER: US 60/404,249
; PRIOR FILING DATE: 2002-08-16
; PRIOR APPLICATION NUMBER: US 60/407,527
; TITLE OF INVENTION: ALPHA GALACTOSIDASE A: RENODELING AND GLYCOCOCONJUGATION OF ALPHA
; FILE REFERENCE: 040853-01-5082
; NUMBER OF SEQ ID NOS: 75
; SOFTWARE: Patentin version 3.2

RESULT 7
US-10-411-026-73
; Sequence 73, Application US/10411026
; Publication No. US2004006391A1
; GENERAL INFORMATION:
; APPLICANT: Neese Technologies, Inc.
; APPLICANT: Derees, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; TITLE OF INVENTION: PROTEIN REMODELING METHODS AND PROTEINS/PEPTIDES PRODUCED BY THE
; FILE REFERENCE: 040853-01-5053
; CURRENT APPLICATION NUMBER: US/10/411,026
; PRIOR FILING DATE: 2003-04-09
; PRIOR APPLICATION NUMBER: US 60/328,523
; PRIOR FILING DATE: 2001-10-0
; PRIOR APPLICATION NUMBER: US 60/344,692
; PRIOR FILING DATE: 2001-10-19
; PRIOR APPLICATION NUMBER: US 60/387,292
; PRIOR FILING DATE: 2002-06-07
; PRIOR APPLICATION NUMBER: US 60/391,777
; PRIOR FILING DATE: 2002-06-25
; PRIOR APPLICATION NUMBER: US 60/396,594
; PRIOR FILING DATE: 2002-07-17
; PRIOR APPLICATION NUMBER: US 60/404,249
; PRIOR FILING DATE: 2002-08-16
; PRIOR APPLICATION NUMBER: US 60/407,527
; TITLE OF INVENTION: ALPHA GALACTOSIDASE A
; FILE REFERENCE: 040853-01-5082
; NUMBER OF SEQ ID NOS: 75
; SOFTWARE: Patentin version 3.2

SEQ ID NO 73
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-411-026-73

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120
Db 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120

QY 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165
Db 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165

RESULT 8
US-10-410-962-73
; Sequence 73, Application US/10410962
; Publication No. US20040077836A1
; GENERAL INFORMATION:
; APPLICANT: Neose Technologies, Inc.
; APPLICANT: DeFrees, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; APPLICANT: Bone, Caryn
; TITLE OF INVENTION: INTERFERON ALPHA: REMODELING AND GLYCOCOCONJUGATION OF INTERFERON
; TITLE OF INVENTION: GLYCOCONJUGATION OF G-CSF
; TITLE OF INVENTION: 040853-01-5054
; CURRENT APPLICATION NUMBER: US/10/410,962
; CURRENT FILING DATE: 2003-04-09
; CURRENT FILING DATE: 2003-04-09
; CURRENT FILING DATE: 2003-04-09
; PRIOR APPLICATION NUMBER: US 60/328,523
; PRIOR FILING DATE: 2002-10-10
; PRIOR FILING DATE: 2002-08-28
; NUMBER OF SEQ ID NOS: 75
; SOFTWARE: Patent version 3.2
; SEQ ID NO 73
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-411-049-73

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120
Db 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120

QY 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165
Db 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165

RESULT 9
US-10-411-049-73
; Sequence 73, Application US/0411049
; Publication No. US20040082026A1
; GENERAL INFORMATION:
; APPLICANT: Neose Technologies, Inc.
; APPLICANT: DeFrees, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; TITLE OF INVENTION: INTERFERON ALPHA: REMODELING AND GLYCOCOCONJUGATION OF INTERFERON
; TITLE OF INVENTION: ALPHA
; TITLE OF INVENTION: 040853-01-5055
; CURRENT APPLICATION NUMBER: US/10/411,049
; CURRENT FILING DATE: 2003-04-09
; CURRENT FILING DATE: 2003-04-09
; CURRENT FILING DATE: 2003-04-09
; PRIOR APPLICATION NUMBER: US 60/391,777
; PRIOR FILING DATE: 2002-06-25
; PRIOR APPLICATION NUMBER: US 60/396,594
; PRIOR FILING DATE: 2002-07-17
; PRIOR APPLICATION NUMBER: US 60/404,249
; PRIOR FILING DATE: 2002-08-16
; PRIOR APPLICATION NUMBER: US 60/407,523
; PRIOR FILING DATE: 2002-08-28
; NUMBER OF SEQ ID NOS: 75
; SOFTWARE: Patent version 3.2
; SEQ ID NO 73
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-411-049-73

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTVPDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120
Db 61 VEWMOGLALISEAVLRGOALLVNSQPWPLQLHVDKAVSGLRSLTTLRGAQKEAIS 120

QY 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165
Db 121 PPDAASAPLRTTADFRKLFRVYSNFRGKLYGEACRTGD 165

RESULT 10
US-10-634-477-1
; Sequence 1, Application US/10634477
; Publication No. US20040110679A1
; GENERAL INFORMATION:
; APPLICANT: Lehmann, Paul
; APPLICANT: Roeddiser, Ralf
; APPLICANT: Walter-Matsui, Ruth
; TITLE OF INVENTION: TREATMENT OF DISTURBANCES OF IRON DISTRIBUTION
; FILE REFERENCE: 21368
; CURRENT APPLICATION NUMBER: US/10/634,477

CURRENT FILING DATE: 2003-08-04
; PRIORITY APPLICATION NUMBER: 02019100.3
; PRIORITY FILING DATE: 2002-08-29
; NUMBER OF SEQ ID NOS: 1
; SOFTWARE: PatentIn Ver. 3.1
; SEQ ID NO: 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-634-477-1

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVILEVRYLLEAKEAENITTGCAHCSLNENITPDTKUNFYAWKRMEMVGQA 60
Db 1 APPRLICDSRVILEVRYLLEAKEAENITTGCAHCSLNENITPDTKUNFYAWKRMEMVGQA 60
QY 61 VEWMOGLALLSEAVLRGQALLVNSQWPPLQLHVDKAVSGRLSLTTLRALGAQEAIIS 120
Db 61 VEWMOGLALLSEAVLRGQALLVNSQWPPLQLHVDKAVSGRLSLTTLRALGAQEAIIS 120

QY 121 PPDAASAPLRTTADTPRKLFVVSNPLRGKLYTGEACTGD 165
Db 121 PPDAASAPLRTTADTPRKLFVVSNPLRGKLYTGEACTGD 165

RESULT 11
US-10-410-930-73
Sequence 73, Application US/10410930
; Publication No. US20040115168A1
; GENERAL INFORMATION:
; APPLICANT: Neose Technologies, Inc.
; APPLICANT: DeFreeze, Shawn
; APPLICANT: Depress, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; APPLICANT: Bowe, Caryn
; TITLE OF INVENTION: POLYCLIC STIMULATING HORMONE: REMODELING AND GLYCOCOCONJUGATION OF
; CURRENT APPLICATION NUMBER: US/10/410,997
; FILE REFERENCE: 040853-01-5059
; PRIOR APPLICATION NUMBER: US 60/328,523
; PRIOR FILING DATE: 2003-04-09
; PRIOR APPLICATION NUMBER: US 60/344,692
; PRIOR FILING DATE: 2003-10-19
; PRIOR APPLICATION NUMBER: US 60/387,292
; PRIOR FILING DATE: 2002-06-07
; PRIOR APPLICATION NUMBER: US 60/391,777
; PRIOR FILING DATE: 2002-06-25
; PRIOR APPLICATION NUMBER: US 60/396,594
; PRIOR FILING DATE: 2002-07-17
; PRIOR APPLICATION NUMBER: US 60/404,249
; PRIOR FILING DATE: 2002-08-16
; PRIOR APPLICATION NUMBER: US 60/407,527
; PRIOR FILING DATE: 2002-08-28
; NUMBER OF SEQ ID NOS: 75
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO: 73
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-410-997-73

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVILEVRYLLEAKEAENITTGCAHCSLNENITPDTKUNFYAWKRMEMVGQA 60
Db 1 APPRLICDSRVILEVRYLLEAKEAENITTGCAHCSLNENITPDTKUNFYAWKRMEMVGQA 60
QY 61 VEWMOGLALLSEAVLRGQALLVNSQWPPLQLHVDKAVSGRLSLTTLRALGAQEAIIS 120
Db 61 VEWMOGLALLSEAVLRGQALLVNSQWPPLQLHVDKAVSGRLSLTTLRALGAQEAIIS 120

QY 121 PPDAASAPLRTTADTPRKLFVVSNPLRGKLYTGEACTGD 165
Db 121 PPDAASAPLRTTADTPRKLFVVSNPLRGKLYTGEACTGD 165

RESULT 13
US-10-411-012-73
; Sequence 73, Application US/10411012
; Publication No. US200413260A1
; GENERAL INFORMATION:
; APPLICANT: Neose Technologies, Inc.

APPLICANT: DeFrees, Shawn ;
 APPLICANT: Zopf, David ;
 APPLICANT: Bayer, Robert ;
 APPLICANT: Hakes, David ;
 APPLICANT: Chen, Xi ;
 APPLICANT: Bone, Caryne ;
 TITLE OF INVENTION: GLYCOPEGYLATION METHODS AND PROTEINS/PEPTIDES PRODUCED BY THE ;
 TITLE OF INVENTION: METHODS ;
 FILE REFERENCE: 040853-01-5051 ;
 CURRENT APPLICATION NUMBER: US 10/411,012 ;
 CURRENT FILING DATE: 2003-04-09 ;
 PRIOR APPLICATION NUMBER: US 60/328,523 ;
 PRIOR FILING DATE: 2001-10-10 ;
 PRIOR APPLICATION NUMBER: US 60/344,692 ;
 PRIOR FILING DATE: 2001-10-19 ;
 PRIOR APPLICATION NUMBER: US 60/387,292 ;
 PRIOR FILING DATE: 2002-06-07 ;
 PRIOR APPLICATION NUMBER: US 60/391,777 ;
 PRIOR FILING DATE: 2002-06-25 ;
 PRIOR APPLICATION NUMBER: US 60/396,594 ;
 PRIOR FILING DATE: 2002-07-17 ;
 PRIOR APPLICATION NUMBER: US 60/404,249 ;
 PRIOR FILING DATE: 2002-08-16 ;
 PRIOR APPLICATION NUMBER: US 60/407,527 ;
 PRIOR FILING DATE: 2002-08-28 ;
 NUMBER OF SEQ ID NOS: 75 ;
 SOFTWARE: PatentIn version 3.2 ;
 SEQ ID NO 73 ;
 LENGTH: 165 ;
 TYPE: PRT ;
 ORGANISM: Homo sapiens ;
 US-10-411-012-73
;
Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
QY 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
Db 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
;
RESULT 14
US-10-410-913-73
;
Sequence 73, Application US/10410913
; Publication No. US20040142856A1
;
GENERAL INFORMATION:
; APPLICANT: Neose Technologies, Inc.
; APPLICANT: DeFrees, Shawn
; APPLICANT: Zopf, David
; APPLICANT: Bayer, Robert
; APPLICANT: Hakes, David
; APPLICANT: Chen, Xi
; APPLICANT: Bone, Caryne
TITLE OF INVENTION: GLYCOCOCONJUGATION METHODS AND PROTEINS/PEPTIDES PRODUCED BY THE ;
TITLE OF INVENTION: METHODS ;
FILE REFERENCE: 040853-01-5081 ;
CURRENT APPLICATION NUMBER: US/10/410,913 ;
CURRENT FILING DATE: 2003-04-09 ;
PRIOR APPLICATION NUMBER: US 60/328,523 ;
PRIOR FILING DATE: 2001-10-10 ;
PRIOR APPLICATION NUMBER: US 60/344,692 ;
PRIOR FILING DATE: 2001-10-19 ;
PRIOR APPLICATION NUMBER: US 60/387,292 ;
PRIOR FILING DATE: 2002-06-07 ;
;
Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
QY 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
Db 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
;
RESULT 15
US-10-780-297-1
;
Sequence 1, Application US/10780297
; Publication No. US20040147431A1
;
GENERAL INFORMATION:
; APPLICANT: Papadimitriou, Apollon
; TITLE OF INVENTION: Erythropoietin Composition
; FILE REFERENCE: 20619 US
; CURRENT APPLICATION NUMBER: US/10/780-297
; CURRENT FILING DATE: 2004-03-17
; PRIOR APPLICATION NUMBER: US/09/853,731
; PRIOR FILING DATE: 2001-05-11
; PRIOR APPLICATION NUMBER: EP/00110355.5
; PRIOR FILING DATE: 2000-05-15
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.0
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-780-297-1
;
Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
QY 1 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENINIVPDTKVNHYAWKRMEVGQOA 60
QY 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
Db 61 VEWMOGLALISEAVLRGQALLVNSSQWPWPLQLAVDKAVSGLSLTLLRALGAQEAKS 120
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
QY 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
Db 121 PPDAASAAPRPTTADTRKLFRVYSNLRGKLYGEACRTGD 165
;

US-10-706-011-1
Sequence 1, Application US/10706701
Publication No. US20040209802A1
GENERAL INFORMATION:
APPLICANT: Lehmann, Paul
APPLICANT: Roeddiger, Ralf
APPLICANT: Walter-Matsui, Ruth
TITLE OF INVENTION: TREATMENT OF DISTURBANCES OF IRON DISTRIBUTION
FILE REFERENCE: 21435
CURRENT APPLICATION NUMBER: US/10/706,701
CURRENT FILING DATE: 2003-11-12
PRIOR APPLICATION NUMBER: 02026342.2
PRIOR FILING DATE: 2002-11-22
NUMBER OF SEQ ID NOS: 1
SOFTWARE: PatentIn Ver. 2.0
SEQ ID NO 1
LENGTH: 165
TYPE: PRT
ORGANISM: Homo sapiens
US-10-706-701-1

Query Match 100.0%; Score 846; DB 4; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
Db 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
QY 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165
Db 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165

RESULT 17

US-10-410-980-73
Sequence 73, Application US/10410980
Publication No. US20050031584A1
GENERAL INFORMATION:
APPLICANT: Neose Technologies, Inc.
APPLICANT: DePree, Shawn
APPLICANT: Zopf, David
APPLICANT: Bayer, Robert
APPLICANT: Hakes, David
APPLICANT: Chen, Xi
APPLICANT: Bowe, Caryn
TITLE OF INVENTION: FACTOR IX: REMODELING AND GLYCOCOCONJUGATION OF FACTOR IX
FILE REFERENCE: 020853-01-5058
CURRENT APPLICATION NUMBER: US/10/410,897
CURRENT FILING DATE: 2003-04-09
PRIOR APPLICATION NUMBER: US 60/328,523
PRIOR FILING DATE: 2001-10-10
PRIOR APPLICATION NUMBER: US 60/344,692
PRIOR FILING DATE: 2001-10-19
PRIOR APPLICATION NUMBER: US 60/387,292
PRIOR FILING DATE: 2002-06-07
PRIOR APPLICATION NUMBER: US 60/391,777
PRIOR FILING DATE: 2002-06-25
PRIOR APPLICATION NUMBER: US 60/396,594
PRIOR FILING DATE: 2002-07-17
PRIOR APPLICATION NUMBER: US 60/404,249
PRIOR FILING DATE: 2002-08-16
PRIOR APPLICATION NUMBER: US 60/407,527
PRIOR FILING DATE: 2002-08-28
NUMBER OF SEQ ID NOS: 75
SOFTWARE: PatentIn version 3.2
SEQ ID NO 73
LENGTH: 165
TYPE: PRT
ORGANISM: Homo sapiens
US-10-410-897-3

Query Match 100.0%; Score 846; DB 5; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
Db 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
QY 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165
Db 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165

RESULT 18

US-10-410-897-73
Sequence 73, Application US/10410987
Publication No. US20050100982A1
GENERAL INFORMATION:
APPLICANT: Neose Technologies, Inc.
APPLICANT: DePree, Shawn
APPLICANT: Zopf, David
APPLICANT: Bayer, Robert
APPLICANT: Hakes, David
APPLICANT: Chen, Xi
APPLICANT: Bowe, Caryn
TITLE OF INVENTION: FACTOR IX: REMODELING AND GLYCOCOCONJUGATION OF FACTOR IX
FILE REFERENCE: 020853-01-5058
CURRENT APPLICATION NUMBER: US/10/410,897
CURRENT FILING DATE: 2003-04-09
PRIOR APPLICATION NUMBER: US 60/328,523
PRIOR FILING DATE: 2001-10-10
PRIOR APPLICATION NUMBER: US 60/344,692
PRIOR FILING DATE: 2001-10-19
PRIOR APPLICATION NUMBER: US 60/387,292
PRIOR FILING DATE: 2002-06-07
PRIOR APPLICATION NUMBER: US 60/391,777
PRIOR FILING DATE: 2002-06-25
PRIOR APPLICATION NUMBER: US 60/396,594
PRIOR FILING DATE: 2002-07-17
PRIOR APPLICATION NUMBER: US 60/404,249
PRIOR FILING DATE: 2002-08-16
PRIOR APPLICATION NUMBER: US 60/407,527
PRIOR FILING DATE: 2002-08-28
NUMBER OF SEQ ID NOS: 75
SOFTWARE: PatentIn version 3.2
SEQ ID NO 73
LENGTH: 165
TYPE: PRT
ORGANISM: Homo sapiens
US-10-410-897-3

Query Match 100.0%; Score 846; DB 5; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
Db 1 APPRLICDSRVRLERYLLEAKEANITGCAERCSLNENITVDTKVNPFYAWKMEVGQA 60
QY 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
Db 61 VEWMOGLLISRAVLRGQALLYNSSQWEPLOLHVDKAVSGRSLTTLRAAGAQEAI 120
QY 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165
Db 121 PPDAASAPLRTTADTPRKLRVYSNPLRGKLYGEACTG 165

RESULT 19
US-11-013-560-1
; Sequence 1, Application US/11013560
; Publication No. US20050181986A1
; GENERAL INFORMATION:
; APPLICANT: WALTER-MATSUI, RUTH
; APPLICANT: RÖDDEGGER, RALF
; APPLICANT: KLIMA, HORST
; TITLE OF INVENTION: METHOD OF TREATING DISTURBANCES OF IRON DISTRIBUTION IN
; FILE REFERENCE: 22351
; CURRENT APPLICATION NUMBER: US/11/013,560
; CURRENT FILING DATE: 2004-12-16
; PRIOR APPLICATION NUMBER: EP 03104932.5
; PRIORITY FILING DATE: 2003-12-19
; NUMBER OF SEQ ID NOS: 4
; SEQ ID NO: 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-013-560-1

Query Match 100.0%; Score 846; DB 6; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; InDelS 0; GapS 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Qy 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Db 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Qy 121 PPDAASAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165
Db 121 PPDAASAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165

RESULT 20
US-09-853-731-2
; Sequence 2, Application US/09853731
; Patent No. US2002003741A1
; GENERAL INFORMATION:

; APPLICANT: Papadimitriou, Apollon
; TITLE OF INVENTION: Erythropoietin Composition
; FILE REFERENCE: 20619 US
; CURRENT APPLICATION NUMBER: US/09/853, 731
; CURRENT FILING DATE: 2001-05-11
; PRIOR APPLICATION NUMBER: EP/00110355.5
; PRIORITY FILING DATE: 2000-05-15
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn version 3.0
; SEQ ID NO: 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-853-731-2

Query Match 100.0%; Score 846; DB 6; Length 165;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; InDelS 0; GapS 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Qy 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Db 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120

RESULT 21
US-10-014-363-2
; Sequence 2, Application US/10014363
; Publication No. US20020211583A1
; GENERAL INFORMATION:
; APPLICANT: BURG, Josef
; APPLICANT: Engel, Alfred
; APPLICANT: Franz, Reinhard
; APPLICANT: Hilger, Bernd
; APPLICANT: Schurig, Hartmut Ernst
; APPLICANT: Fischer, Wilhelm
; TITLE OF INVENTION: Erythropoietin Conjugates
; FILE REFERENCE: Case 2005
; CURRENT APPLICATION NUMBER: US/10/014,363
; CURRENT FILING DATE: 2001-12-11
; NUMBER OF SEQ ID NOS: 5
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO: 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-014-363-2

Query Match 100.0%; Score 846; DB 4; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; InDelS 0; GapS 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Qy 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Db 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Qy 121 PPDAASAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165
Db 121 PPDAASAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165

RESULT 22
US-10-241-356-2
; Sequence 2, Application US/10241356
; Publication No. US2003007753A1
; GENERAL INFORMATION:
; APPLICANT: TISCHER, WILHELM
; TITLE OF INVENTION: DIGLYCOSYLATED ERYTHROPOETIN
; FILE REFERENCE: 20971
; CURRENT APPLICATION NUMBER: US/10/241,356
; CURRENT FILING DATE: 2002-09-11
; PRIOR APPLICATION NUMBER: EP 0122555.4
; PRIORITY FILING DATE: 2001-09-25
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO: 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-241-356-2

Query Match 100.0%; Score 846; DB 4; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; InDelS 0; GapS 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLENENTVDPDKVNHYAWKRMVEGQQA 60
Qy 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120
Db 61 VEWVGGLALLSEAVLRQQLIVNSQQPWEPLQLRHDKAVSGLRSLTTLRALGAQEKAIS 120

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; SOFTWARE: PatentIn Ver. 2.0
; SEQ_ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-400-377-2

QY 1 APPRILICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60
Db 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120

RESULT 23
US-10-293-551-2
; Sequence 2, Application US/10293551
; Publication No. US20030120045A1
; GENERAL INFORMATION:
; APPLICANT: Bailon, Pascal
; TITLE OF INVENTION: BRYTHRPOPOETIN CONJUGATES
; CURRENT APPLICATION NUMBER: US/10/293,551
; CURRENT FILING DATE: 2002-11-14
; PRIOR APPLICATION NUMBER: US/09/604,938
; PRIOR FILING DATE: 2000-06-27
; PRIOR APPLICATION NUMBER: 60/166,151
; PRIOR FILING DATE: 1999-11-17
; PRIOR APPLICATION NUMBER: 60/151,548
; PRIOR FILING DATE: 1999-08-13
; PRIOR APPLICATION NUMBER: 60/150,225
; PRIOR FILING DATE: 1999-08-23
; PRIOR APPLICATION NUMBER: 60/142,254
; PRIOR FILING DATE: 1999-07-02
; NUMBER OF SEQ ID NOS: 3
; SOFTWARE: Patentin Ver. 2.1
SEQ ID NO 2
LENGTH: 166
TYPE: PRT
ORGANISM: Homo sapiens
US-10-293-551-2

Query Match 100.0%; Score 846; DB 4; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRILICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60
Db 1 APPRLICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60

QY 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
Db 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120

RESULT 25
US-10-400-708-2
; Sequence 2, Application US/10400708
; Publication No. US20030166865A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; FILE REFERENCE: 4152-1-PUS
; CURRENT APPLICATION NUMBER: US/10/400,708
; CURRENT FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: US/09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: 60/052,516
; NUMBER OF SEQ ID NOS: 41
; SOFTWARE: Patentin Ver. 2.0
SEQ ID NO 2
LENGTH: 166
TYPE: PRT
ORGANISM: Homo sapiens
US-10-400-708-2

Query Match 100.0%; Score 846; DB 4; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRILICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60
Db 1 APPRLICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60

QY 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
Db 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120

QY 121 PPDRASAAPLRTTADFRKLFRVYSNFLRGKLYGEACRTGD 165
Db 121 PPDRASAAPLRTTADFRKLFRVYSNFLRGKLYGEACRTGD 165
121 PPDRASAAPLRTTADFRKLFRVYSNFLRGKLYGEACRTGD 165

RESULT 24
US-10-400-377-2
; Sequence 2, Application US/10400377
; Publication No. US030162949A1
; GENERAL INFORMATION:
; APPLICANT: Bolder Biotechnology, Inc.
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; CURRENT APPLICATION NUMBER: US/10/400,377
; CURRENT FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: US/09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; NUMBER OF SEQ ID NOS: 41

QY 1 APPRILICDSRVLYLEAKEAENITTCGAECISLNENITVPDKVNFTYAWKMEVGQQA 60
Db 61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120
61 VEWQGLALLSEAVLRGQALVNSSQPPEPLQHVDKAVSGLRSITTLRALGQKEAIS 120

RESULT 26
US-10-298-148-2
; Sequence 2, Application US/10298148
; Publication No. US20030171284A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; APPLICANT: Bolder Biotechnology, Inc.
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; FILE REFERENCE: 4152-1-PUS

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; CURRENT APPLICATION NUMBER: US/10/298,148 ; Sequence 1, Application US/10467115
; CURRENT FILING DATE: 2002-11-15 ; Publication No. US20040063917A1
; PRIORITY NUMBER: US/09/462,941 ; GENERAL INFORMATION:
; PRIORITY NUMBER: 60/052,516 ; APPLICANT: Carr, Francis J.
; PRIORITY FILING DATE: 2000-01-14 ; APPLICANT: Carter, Graham
; SEQ ID NO: 2 ; APPLICANT: Jones, Tim
; LENGTH: 166 ; APPLICANT: Williams, Stephen
; TYPE: PRT ; TITLE OF INVENTION: MODIFIED ERYTHROPOEITIN (EPO) WITH
; ORGANISM: Homo sapiens ; TITLE OF INVENTION: REDUCED IMMUNOGENICITY
; US-10-298-148-2 ; FILE REFERENCE: MER-114
; ; CURRENT APPLICATION NUMBER: US/10/467,115
; ; CURRENT FILING DATE: 2003-08-05
; ; PRIORITY NUMBER: 01102615.0
; ; PRIORITY FILING DATE: 2001-02-06
; ; PRIORITY APPLICATION NUMBER: 01103954.2
; ; PRIORITY FILING DATE: 2001-02-19
; ; PRIORITY APPLICATION NUMBER: PCT/EP02/01174
; ; PRIORITY FILING DATE: 2002-02-05
; ; NUMBER OF SEQ ID NOS: 72
; ; SOFTWARE: FastSEQ for Windows Version 4.0
; ; SEQ ID NO: 1
; ; LENGTH: 166
; ; TYPE: PRT
; ; ORGANISM: Homo sapien
; ; US-10-467-115-1
; Query Match 100.0%; Score 846; DB 4; Length 166;
; Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDKYKVNFKWKRMEVGQA 60
; Db 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDKYKVNFKWKRMEVGQA 60
; QY 61 VEWQGALLSEAVLRGQALLVNSQWPBPLQHVDKAVSGLRSLTTLRALGAQEAKTS 120
; Db 61 VEWQGALLSEAVLRGQALLVNSQWPBPLQHVDKAVSGLRSLTTLRALGAQEAKTS 120
; QY 121 PPDAASAAPLRTTADTRKLFRVYNSFLRGKLYGEACRTGD 165
; Db 121 PPDAASAAPLRTTADTRKLFRVYNSFLRGKLYGEACRTGD 165
; RESULT 27
; ; Sequence 227, Application US/10360101
; ; Publication No. US20040009550A1
; ; GENERAL INFORMATION:
; ; APPLICANT: Moll, Gert N.
; ; APPLICANT: Leenhouts, Cornelis J.
; ; TITLE OF INVENTION: Export and modification of (poly)peptide in the lantibiotic way
; ; FILE REFERENCE: 2183-5673
; ; CURRENT APPLICATION NUMBER: US/10/360,101
; ; CURRENT FILING DATE: 2003-02-07
; ; PRIORITY APPLICATION NUMBER: EP 02077060.8
; ; PRIORITY FILING DATE: 2002-05-24
; ; NUMBER OF SEQ ID NOS: 309
; ; SEQ ID NO: 227
; ; LENGTH: 166
; ; TYPE: PRT
; ; ORGANISM: Artificial Sequence
; ; FEATURE:
; ; OTHER INFORMATION: sequence of erythropoietin
; ; US-10-360-101-227
; Query Match 100.0%; Score 846; DB 4; Length 166;
; Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDKYKVNFKWKRMEVGQA 60
; Db 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENNTVPDKYKVNFKWKRMEVGQA 60
; QY 61 VEWQGALLSEAVLRGQALLVNSQWPBPLQHVDKAVSGLRSLTTLRALGAQEAKTS 120
; Db 61 VEWQGALLSEAVLRGQALLVNSQWPBPLQHVDKAVSGLRSLTTLRALGAQEAKTS 120
; QY 121 PPDAASAAPLRTTADTRKLFRVYNSFLRGKLYGEACRTGD 165
; Db 121 PPDAASAAPLRTTADTRKLFRVYNSFLRGKLYGEACRTGD 165
; RESULT 29
; ; Sequence 201, Application US/10658834A
; ; Publication No. US20040132977A1
; ; GENERAL INFORMATION:
; ; APPLICANT: Gantier, Rene
; ; APPLICANT: Guyon, Thierry
; ; APPLICANT: Drittanti, Lila
; ; APPLICANT: Vega, Manuel
; ; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding N
; ; TITLE OF INVENTION: Molecules and Related Applications
; ; FILE REFERENCE: 3751-922
; ; CURRENT APPLICATION NUMBER: US/10/658,834A
; ; CURRENT FILING DATE: 2003-03-08
; ; PRIORITY NUMBER: 60/457,135
; ; PRIORITY FILING DATE: 2003-03-21
; ; PRIORITY APPLICATION NUMBER: 3751-922
; ; PRIORITY FILING DATE: 2002-09-09
; ; NUMBER OF SEQ ID NOS: 1306
; ; SOFTWARE: FastSEQ for Windows Version 4.0
; ; SEQ ID NO: 201
; ; LENGTH: 166
; ; TYPE: PRT
; ; ORGANISM: Homo sapiens
; ; PUBLICATION INFORMATION:
; ; DATABASE ACCESSION NUMBER: Genbank AAA52400
; ; DATABASE ENTRY DATE: 1994-11-08
; ; US-10-658-834-201
; RESULT 28
; ; US-10-467-115-1

```


APPLICANT: Carter, Graham
 APPLICANT: Jones, Tim
 APPLICANT: Williams, Stephen
 APPLICANT: Hamilton, Anita
 TITLE OF INVENTION: METHOD FOR IDENTIFICATION OF T-CELL
 FILE REFERENCE: MER-117
 CURRENT APPLICATION NUMBER: US/10/468,496
 PRIOR APPLICATION NUMBER: 01103954.2
 PRIOR FILING DATE: 2001-07-19
 PRIOR APPLICATION NUMBER: 01105777.5
 PRIOR FILING DATE: 2001-01-08
 PRIOR APPLICATION NUMBER: 01106538.0
 PRIOR FILING DATE: 2001-03-15
 PRIOR APPLICATION NUMBER: 01106536.4
 PRIOR FILING DATE: 2001-03-15
 PRIOR APPLICATION NUMBER: 01107012.5
 PRIOR FILING DATE: 2001-03-20
 PRIOR APPLICATION NUMBER: 01106899.6
 PRIOR FILING DATE: 2001-03-20
 NUMBER OF SEQ ID NOS: 2036
 SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 133
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo Sapiens
; US-10-468-496-133

Query Match 100.0%; Score 846; DB 4; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60
 Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60

Qy 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120
 Db 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120

Qy 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165

RESULT 34
 Sequence 2, Application US/10773-654-2
 Publication No. US20040214287A1
 GENERAL INFORMATION:
 APPLICANT: Cox III, George N
 APPLICANT: Bolder Biotechnology, Inc.
 TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
 FILE REFERENCE: 4152-1-PUS
 CURRENT APPLICATION NUMBER: US/10/866,540
 PRIOR APPLICATION NUMBER: US/10/460,377
 PRIOR FILING DATE: 2003-03-26
 PRIOR APPLICATION NUMBER: US/09/462,941
 PRIOR FILING DATE: 2000-01-14
 PRIOR APPLICATION NUMBER: 6/0/052,516
 PRIOR FILING DATE: 1997-07-14
 NUMBER OF SEQ ID NOS: 41
 SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-866-540-2

Query Match 100.0%; Score 846; DB 5; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60
 Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60

Qy 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120
 Db 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120

Qy 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165

RESULT 35
 Sequence 2, Application US/10866-540-2
 Publication No. US20040230040A1
 GENERAL INFORMATION:
 APPLICANT: Cox III, George N
 APPLICANT: Bolder Biotechnology, Inc.
 TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
 FILE REFERENCE: 4152-1-PUS
 CURRENT APPLICATION NUMBER: US/10/866,540
 PRIOR APPLICATION NUMBER: US/10/460,377
 PRIOR FILING DATE: 2003-03-26
 PRIOR APPLICATION NUMBER: US/09/462,941
 PRIOR FILING DATE: 2000-01-14
 PRIOR APPLICATION NUMBER: 6/0/052,516
 PRIOR FILING DATE: 1997-07-14
 NUMBER OF SEQ ID NOS: 41
 SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-866-540-2

Query Match 100.0%; Score 846; DB 5; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.5e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60
 Db 1 APPRLICDSVRLERYLLEAKEAENITGCAEHCSLENNTIVPDTKVNFKYAWKRMVGQAA 60

Qy 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120
 Db 61 VEVWQGLALISEAVLRGQALLVNSQWPWPLQLHYDKAVSGLSRSLTLLRALGAQEATS 120

Qy 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165
 Db 121 PPDASAAPLRTTADTRKLFRVYSNFLRGKLUYGEACRTGD 165

RESULT 36
 Sequence 2, Application US/10856-219-2
 Publication No. US20040255269A1
 GENERAL INFORMATION:
 APPLICANT: Cox III, George N
 APPLICANT: Bolder Biotechnology, Inc.
 TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
 FILE REFERENCE: 4152-1-PUS
 CURRENT APPLICATION NUMBER: US/10/856,219
 PRIOR APPLICATION NUMBER: US/10/400,377
 PRIOR FILING DATE: 2003-03-26
 PRIOR APPLICATION NUMBER: US/09/462,941
 PRIOR FILING DATE: 2000-01-14

Query Match 100.0%; Score 846; DB 4; Length 166;

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; PRIORITY APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; NUMBER OF SEQ ID NOS: 41
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-856-219-2

Query Match Similarity 100.0%; Score 846; DB 5; Length 166;
Best Local Similarity 100.0%; Pred. No. 1 5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Sequence 2, Application US/10685288
; Publication No. US20050058621A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; TITER OF INVENTION: Thereof
; FILE REFERENCE: 4152-1-PUS-8
; CURRENT APPLICATION NUMBER: US/10/685, 288
; CURRENT FILING DATE: 2003-10-13
; PRIOR APPLICATION NUMBER: 60/418,106
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/418,105
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 10/400,377
; PRIOR FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: 09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: PCT/US98/14497
; PRIOR FILING DATE: 1998-07-13
; PRIOR APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; PRIOR FILING DATE: 2002-11-15
; PRIOR APPLICATION NUMBER: 60/418,040
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/332,285
; PRIOR FILING DATE: 2001-11-15
; PRIOR APPLICATION NUMBER: 09/889,273
; PRIOR FILING DATE: 2001-07-13
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 41
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-685-288-2

Query Match Similarity 100.0%; Score 846; DB 5; Length 166;
Best Local Similarity 100.0%; Pred. No. 1 5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Sequence 2, Application US/10866580
; Publication No. US20050096461A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; CURRENT APPLICATION NUMBER: US/10/866, 580
; CURRENT FILING DATE: 2004-06-10
; PRIOR APPLICATION NUMBER: US/10/400,377
; PRIOR FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: US/09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-866-580-2

Query Match Similarity 100.0%; Score 846; DB 5; Length 166;
Best Local Similarity 100.0%; Pred. No. 1 5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Sequence 2, Application US/10866580
; Publication No. US20050096461A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; CURRENT APPLICATION NUMBER: US/10/866, 580
; CURRENT FILING DATE: 2004-06-10
; PRIOR APPLICATION NUMBER: US/10/400,377
; PRIOR FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: US/09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 2
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-10-866-580-2

Query Match Similarity 100.0%; Score 846; DB 5; Length 166;
Best Local Similarity 100.0%; Pred. No. 1 5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Sequence 2, Application US/10773530
; Publication No. US20050107591A1
; GENERAL INFORMATION:
; APPLICANT: Cox III, George N
; TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
; CURRENT APPLICATION NUMBER: US/10/773, 530
; CURRENT FILING DATE: 2004-02-05
; PRIOR APPLICATION NUMBER: US/10/400,377
; PRIOR FILING DATE: 2003-03-26
; PRIOR APPLICATION NUMBER: US/09/462,941
; PRIOR FILING DATE: 2000-01-14
; PRIOR APPLICATION NUMBER: 60/052,516
; PRIOR FILING DATE: 1997-07-14
; SOFTWARE: PatentIn Ver. 2.0
; NUMBER OF SEQ ID NOS: 41

```


RESULT 43
US-10-014-363-4
Sequence 4, Application US/10014363
Publication No. US20020115833A1
GENERAL INFORMATION:
APPLICANT: Burg, Josef
APPLICANT: Engel, Alfred
APPLICANT: Franz, Reinhard
APPLICANT: Hilger, Bernd
APPLICANT: Schurig, Hartmut Ernst
APPLICANT: Tischer, Wilhelm
APPLICANT: Wozny, Manfred
TITLE OF INVENTION: Erythropoietin Conjugates
FILE REFERENCE: Case 20805
CURRENT APPLICATION NUMBER: US/10/014, 363
CURRENT FILING DATE: 2001-12-11
NUMBER OF SEQ ID NOS: 5
SOFTWARE: PatentIn version 3.1
SEQ ID NO 4
LENGTH: 169
TYPE: PRT
ORGANISM: CHO/dhfr-
US-10-014-363-4

Query Match 100.0%; Score 846; DB 4; Length 169;
Best Local Similarity 100.0%; Pred. No. 1. 5e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Del 0; Gap 0;

QY 1 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 60
Db 4 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 63

QY 61 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 120
Db 64 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 123

QY 121 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 165
Db 124 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 168

RESULT 44
US-10-014-363-3
Sequence 3, Application US/10014363
Publication No. US20020115833A1
GENERAL INFORMATION:
APPLICANT: Burg, Josef
APPLICANT: Engel, Alfred
APPLICANT: Franz, Reinhard
APPLICANT: Hilger, Bernd
APPLICANT: Schurig, Hartmut Ernst
APPLICANT: Tischer, Wilhelm
APPLICANT: Wozny, Manfred
TITLE OF INVENTION: Erythropoietin Conjugates
FILE REFERENCE: Case 20805
CURRENT APPLICATION NUMBER: US/10/014, 363
NUMBER OF SEQ ID NOS: 5
SOFTWARE: PatentIn version 3.1
SEQ ID NO 5
LENGTH: 174
TYPE: PRT
ORGANISM: CHO/dhfr-
US-10-014-363-5

Query Match 100.0%; Score 846; DB 4; Length 174;
Best Local Similarity 100.0%; Pred. No. 1. 6e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Del 0; Gap 0;

QY 1 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 60
Db 9 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 68

QY 61 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 120
Db 69 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 128

QY 121 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 165
Db 129 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 173

RESULT 45
US-10-014-363-5
Sequence 5, Application US/10014363
Publication No. US20020115833A1
GENERAL INFORMATION:
APPLICANT: Franz, Reinhard
APPLICANT: Hilger, Bernd
APPLICANT: Schurig, Hartmut Ernst
APPLICANT: Tischer, Wilhelm
APPLICANT: Wozny, Manfred
TITLE OF INVENTION: Erythropoietin Conjugates
FILE REFERENCE: Case 20805
CURRENT APPLICATION NUMBER: US/10/014, 363
NUMBER OF SEQ ID NOS: 5
SOFTWARE: PatentIn version 3.1
SEQ ID NO 5
LENGTH: 174
TYPE: PRT
ORGANISM: CHO/dhfr-
US-10-014-363-5

Query Match 100.0%; Score 846; DB 4; Length 174;
Best Local Similarity 100.0%; Pred. No. 1. 6e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Del 0; Gap 0;

QY 1 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 60
Db 9 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 68

QY 61 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 120
Db 69 VEWQGLALLSERVLRGQALLVNSQWPQLQVNDKAVSGIERSLTILRAAGQEAIS 128

QY 121 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 165
Db 129 PPDASAAPLRTTADTPRKLRVYSNPLRGKLYGEACTGQ 173

RESULT 46
US-10-775-204-593
Sequence 593, Application US/10775204
Publication No. US20050186664A1
GENERAL INFORMATION:
APPLICANT: Rauen, Craig A.
APPLICANT: Baseline, William A.
APPLICANT: Balance, David J.
APPLICANT: Turner, Andrew J.
TITLE OF INVENTION: Alumin Fusion Proteins
FILE REFERENCE: P2564
CURRENT APPLICATION NUMBER: US/10/775, 204
CURRENT FILING DATE: 2004-02-11
PRIORITY APPLICATION NUMBER: 60/341, 811
PRIORITY FILING DATE: 2003-12-21
PRIORITY APPLICATION NUMBER: 60/360, 000
PRIORITY FILING DATE: 2003-02-28
PRIORITY APPLICATION NUMBER: 60/378, 950
PRIORITY FILING DATE: 2002-05-10
PRIORITY APPLICATION NUMBER: 60/398, 008
PRIORITY FILING DATE: 2002-07-24

Query Match 100.0%; Score 846; DB 4; Length 174;
Best Local Similarity 100.0%; Pred. No. 1. 6e-85; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Del 0; Gap 0;

QY 1 APPLICODSRVRLERYLKEAKENITTCGAECNSLENITVDPDKVNYAWKMEVGQA 60

PRIOR FILING DATE: 2002-05-10
 PRIOR APPLICATION NUMBER: 60/498,008
 PRIOR FILING DATE: 2002-07-24
 PRIOR APPLICATION NUMBER: 60/411,355
 PRIOR FILING DATE: 2002-09-18
 PRIOR APPLICATION NUMBER: 60/414,984
 PRIOR APPLICATION NUMBER: 60/417,611
 PRIOR APPLICATION NUMBER: 60/420,246
 PRIOR FILING DATE: 2002-10-23
 PRIOR APPLICATION NUMBER: 60/423,623
 PRIOR FILING DATE: 2002-11-05
 PRIOR APPLICATION NUMBER: 60/351,360

PRIOR APPLICATION NUMBER: 60/351,360
 PRIOR FILING DATE: 2002-01-28
 Remaining prior Application data removed - See File Wrapper or PALM.
 NUMBER OF SEQ ID NOS: 2222
 SOFTWARE: Patentin Ver. 2.0
 SEQ ID NO 1828
 LENGTH: 192
 TYPE: PRT
 ORGANISM: Homo sapiens

US-10-775-204-1691

Query Match 100.0%; Score 846; DB 5; Length 192;
 Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0;

b 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENTVDPDKVNFTAWKRMEVGQOA 60
 28 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENTVDPDKVNFTAWKRMEVGQOA 87

y 61 VEVNQGLALLSEAVLRGQALLVNSQWPBLQHVDKAVSGRLSLTLLRALGAQEAKS 120
 88 VEVNQGLALLSEAVLRGQALLVNSQWPBLQHVDKAVSGRLSLTLLRALGAQEAKS 147

Y 121 PPDAASAAPLRTTADTPRKLFRLFRVYSNPLRGKLYTGEACRTGD 165
 Y 148 PPDAASAAPLRTTADTPRKLFRLFRVYSNPLRGKLYTGEACRTGD 192

RESULT 52

S-10-775-204-1828

Publication 1828, Application US/10775204

GENERAL INFORMATION:

APPLICANT: Rosen, Craig A.

APPLICANT: Haseltine, William A.

APPLICANT: Balance, David J.

APPLICANT: Turner, Andrew J.

TITLE OF INVENTION: Albumin Fusion Proteins

FILE REFERENCE: PF564

CURRENT APPLICATION NUMBER: US/10/775,204

PRIOR FILING DATE: 2004-02-11

PRIOR APPLICATION NUMBER: 60/341,811

PRIOR FILING DATE: 2001-12-21

PRIOR APPLICATION NUMBER: 60/360,000

PRIOR FILING DATE: 2002-02-28

PRIOR APPLICATION NUMBER: 60/378,950

PRIOR FILING DATE: 2002-05-10

PRIOR APPLICATION NUMBER: 60/398,008

PRIOR FILING DATE: 2002-07-24

PRIOR APPLICATION NUMBER: 60/411,355

PRIOR FILING DATE: 2002-09-18

PRIOR APPLICATION NUMBER: 60/414,984

PRIOR FILING DATE: 2002-10-02

PRIOR APPLICATION NUMBER: 60/417,611

PRIOR FILING DATE: 2002-10-11

PRIOR APPLICATION NUMBER: 60/411,355

PRIOR FILING DATE: 2002-09-18

PRIOR APPLICATION NUMBER: 60/414,984

PRIOR FILING DATE: 2002-10-02

PRIOR APPLICATION NUMBER: 60/398,008

PRIOR FILING DATE: 2002-07-24

PRIOR APPLICATION NUMBER: 60/420,246

PRIOR FILING DATE: 2002-10-23

PRIOR APPLICATION NUMBER: 60/423,623

PRIOR FILING DATE: 2002-11-05

PRIOR APPLICATION NUMBER: 60/351,360

PRIOR FILING DATE: 2002-01-28

Remaining Prior Application data removed - See File Wrapper or PALM.

NUMBER OF SEQ ID NOS: 2222
 SOFTWARE: Patentin Ver. 2.0
 SEQ ID NO 1829
 LENGTH: 192
 TYPE: PRT
 ORGANISM: Homo sapiens

US-10-775-204-1828

Query Match 100.0%; Score 846; DB 5; Length 192;
 Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Mismatches 0; Indels 0; Gaps 0;

Db 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENTVDPDKVNFTAWKRMEVGQOA 60
 28 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENTVDPDKVNFTAWKRMEVGQOA 87

Qy 61 VEVNQGLALLSEAVLRGQALLVNSQWPBLQHVDKAVSGRLSLTLLRALGAQEAKS 120
 88 VEVNQGLALLSEAVLRGQALLVNSQWPBLQHVDKAVSGRLSLTLLRALGAQEAKS 147

Db 121 PPDAASAAPLRTTADTPRKLFRLFRVYSNPLRGKLYTGEACRTGD 165
 Db 148 PPDAASAAPLRTTADTPRKLFRLFRVYSNPLRGKLYTGEACRTGD 192

RESULT 53

US-10-775-204-1829

; Sequence 1829, Application US/10775204

; Publication No. US2005018664A1

; GENERAL INFORMATION:

; APPLICANT: Rosen, Craig A.

; APPLICANT: Haseltine, William A.

; APPLICANT: Balance, David J.

; APPLICANT: Turner, Andrew J.

; TITLE OF INVENTION: Albumin Fusion Proteins

; FILE REFERENCE: PF564

; CURRENT APPLICATION NUMBER: US/10/775,204

; CURRENT FILING DATE: 2004-02-11

; PRIOR FILING DATE: 2001-12-21

; PRIOR APPLICATION NUMBER: 60/341,811

; PRIOR FILING DATE: 2004-02-11

; PRIOR APPLICATION NUMBER: 60/360,000

; PRIOR FILING DATE: 2002-02-28

; PRIOR APPLICATION NUMBER: 60/378,950

; PRIOR FILING DATE: 2002-05-10

; PRIOR APPLICATION NUMBER: 60/398,008

; PRIOR FILING DATE: 2002-07-24

; PRIOR APPLICATION NUMBER: 60/411,355

; PRIOR FILING DATE: 2002-09-18

; PRIOR APPLICATION NUMBER: 60/414,984

; PRIOR FILING DATE: 2002-10-02

; PRIOR APPLICATION NUMBER: 60/417,611

; PRIOR FILING DATE: 2002-10-11

; PRIOR APPLICATION NUMBER: 60/420,246

; PRIOR FILING DATE: 2002-10-23

; PRIOR APPLICATION NUMBER: 60/423,623

; PRIOR FILING DATE: 2002-11-05

; PRIOR APPLICATION NUMBER: 60/351,360

; PRIOR FILING DATE: 2002-01-28

; Remaining Prior Application data removed - See File Wrapper or PALM.

; NUMBER OF SEQ ID NOS: 2222
 ; SOFTWARE: Patentin Ver. 2.0
 ; SEQ ID NO 1829
 ; LENGTH: 192
 ; TYPE: PRT
 ; ORGANISM: Homo sapiens

US-10-775-204-1829

Query Match 100.0%; Score 846; DB 5; Length 192;
 Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 60 US-09-813-775C-4
Db 28 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 87 ; Sequence 4, Application US/09813775C
; Publication No. US20030054494A1
; GENERAL INFORMATION:
QY 61 VEVWQGLALLSEAVLRGQALLVNSQPEPLQLHVDKAVSGRSLSLTLLRAIGAOKEAIIS 120 ; APPLICANT: Desauvage, Frederick
; APPLICANT: Henner, Dennis J.
Db 88 VEVWQGLALLSEAVLRGQALLVNSQPEPLQLHVDKAVSGRSLSLTLLRAIGAOKEAIIS 147 ; APPLICANT: Baseltine, William A.
; TITLE OF INVENTION: No. US20030054494A1
; TITLE OF INVENTION: Polypeptides and nucleic acids encoding the same
QY 121 PPDAAASAPLRTTADTRKLFRVYSNLRGKLUKYGEACRTGD 165 FILE REFERENCE: GENENT_057622
Db 148 PPDAAASAPLRTTADTRKLFRVYSNLRGKLUKYGEACRTGD 192 CURRENT APPLICATION NUMBER: US/09-813,775C
; CURRENT FILING DATE: 1999-05-07
; PRIOR APPLICATION NUMBER: US 09/307307
; PRIOR FILING DATE: 1999-05-07
; PRIOR APPLICATION NUMBER: US 09/552265
; PRIOR FILING DATE: 2000-04-19
; NUMBER OF SEQ ID NOS: 52 SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 4
; LENGTH: 193
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-813-775C-4

RESULT 54
US-10-775-204-1830
; Sequence 1830, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Baseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: PP564
; CURRENT APPLICATION NUMBER: US/10/775, 204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341,811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/336,000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378,950
; PRIOR FILING DATE: 2002-05-10
; PRIOR APPLICATION NUMBER: 60/398,008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411,355
; PRIOR FILING DATE: 2002-09-18
; PRIOR APPLICATION NUMBER: 60/414,984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/417,611
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/420,246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423,623
; PRIOR FILING DATE: 2002-11-05
; PRIOR APPLICATION NUMBER: 60/351,360
; PRIOR FILING DATE: 2002-01-28
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO: 1830
; LENGTH: 192
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-1830

RESULT 55
US-10-113-824-2
; Sequence 2, Application US/10113824
; Publication No. US20030050269A1
; GENERAL INFORMATION:
; APPLICANT: Bacary, Jean-Louis
; TITLE OF INVENTION: NEW POLYNUCLEOTIDES AND POLYPEPTIDES OF THE ERYTHROPOIETIN GENE
; FILE REFERENCE: 02134910037
; CURRENT APPLICATION NUMBER: US/10/113, 824
; CURRENT FILING DATE: 2002-03-29
; PRIOR APPLICATION NUMBER: FR 0104603
; PRIOR FILING DATE: 2001-04-04
; PRIOR APPLICATION NUMBER: US 60/343163
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: US 60/345,440
; PRIOR FILING DATE: 2002-01-04
; PRIOR APPLICATION NUMBER: US 60/358,598
; NUMBER OF SEQ ID NOS: 22
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO: 2
; LENGTH: 193
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-113-824-2

Query Match 100 %; Score 846; DB 5; Length 192;
Best Local Similarity 100.0%; Pred. No. 1.8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 60
Db 28 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 87
; Query Match 100 %; Score 846; DB 4; Length 193;
Best Local Similarity 100.0%; Pred. No. 1.8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 61 VEVWQGLALLSEAVLRGQALLVNSQPEPLQLHVDKAVSGRSLSLTLLRAIGAOKEAIIS 120
Db 88 VEVWQGLALLSEAVLRGQALLVNSQPEPLQLHVDKAVSGRSLSLTLLRAIGAOKEAIIS 147
; Query Match 100 %; Score 846; DB 4; Length 193;
Best Local Similarity 100.0%; Pred. No. 1.8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 60
Db 28 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVPTDKUNFYAWKRMVQQA 87

RESULT 55

RESULT 57

US-10-612-665-10

; Sequence 10, Application US/10612665

; Publication No. US20040122216A1

; GENERAL INFORMATION:

QY 121 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 165

Db 148 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 192

QY ; APPLICANT: Nielsen, J.

Db ; APPLICANT: Pedersen, L.

QY ; APPLICANT: Christensen, S.

Db ; APPLICANT: Sager, T.

QY ; APPLICANT: Brines, M.

Db ; APPLICANT: Leist, M.

QY ; APPLICANT: Geist, M.

Db ; APPLICANT: Kallunki, P.

QY ; APPLICANT: Nielsen, J.

Db ; APPLICANT: Pedersen, J.

QY ; APPLICANT: Gerwien, J.

Db ; APPLICANT: Bay, K.

QY ; APPLICANT: Pedersen, L.

Db ; APPLICANT: Leist, M.

QY ; APPLICANT: Geist, M.

Db ; APPLICANT: Kallunki, P.

QY ; APPLICANT: Christensen, S.

Db ; APPLICANT: Sager, T.

QY ; APPLICANT: Brines, M.

Db ; APPLICANT: Cerami, A.

QY ; APPLICANT: Cerami, C.

Db ; TITLE OF INVENTION: RECOMBINANT TISSUE PROTECTIVE CYTOKINES AND ENCODING NUCLEIC ACIDS THEREOF FOR PROTECTION, RESTORATION, AND ENHANCEMENT OF TISSUES AND ORGANS

QY ; TITLE OF INVENTION: RECOMBINANT TISSUE PROTECTIVE CYTOKINES AND ENCODING NUCLEIC ACIDS THEREOF FOR PROTECTION, RESTORATION, AND ENHANCEMENT OF TISSUES AND ORGANS

Db ; FILE REFERENCE: 10165-022-999

QY ; CURRENT FILING DATE: 2003-07-01

Db ; PRIOR APPLICATION NUMBER: 60/392,455

QY ; PRIOR FILING DATE: 2002-07-01

Db ; PRIOR APPLICATION NUMBER: 60/393,423

QY ; PRIOR FILING DATE: 2002-07-03

Db ; NUMBER OF SEQ ID NOS: 212

QY ; SEQ ID NO: 22

Db ; SOFTWARE: PatentIn version 3.2

QY ; LENGTH: 193

Db ; TYPE: PRT

QY ; ORGANISM: Artificial

Db ; PARENT: Cerami, C.

QY ; OTHER INFORMATION: Description of Artificial Sequence: mutein

US-10-612-665-22

Query Match 100.0%; Score 846; DB 4; Length 193;

Best Local Similarity 100.0%; Pred. No. 1.8e-85; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRICDSVRLEVYLLEAKEAENTITGCACHECSLNENTVPDTKVNFYAWKMEVGQOA 60

Db 28 APPRICDSVRLEVYLLEAKEAENTITGCACHECSLNENTVPDTKVNFYAWKMEVGQOA 87

QY 61 VEWQGLALISEAUTRGQLVNSQPWPFLQIHDVKAVSGLSLTTLRALGAQEAKIS 120

Db 88 VEWQGLALISEAUTRGQLVNSQPWPFLQIHDVKAVSGLSLTTLRALGAQEAKIS 147

QY 121 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 165

Db 148 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 192

RESULT 59

US-10-612-665-112

; Sequence 12, Application US/10612665

; Publication No. US20040122216A1

; GENERAL INFORMATION:

QY 1 APPRICDSVRLEVYLLEAKEAENTITGCACHECSLNENTVPDTKVNFYAWKMEVGQOA 60

Db 28 APPRICDSVRLEVYLLEAKEAENTITGCACHECSLNENTVPDTKVNFYAWKMEVGQOA 87

QY 61 VEWQGLALISEAUTRGQLVNSQPWPFLQIHDVKAVSGLSLTTLRALGAQEAKIS 120

Db 88 VEWQGLALISEAUTRGQLVNSQPWPFLQIHDVKAVSGLSLTTLRALGAQEAKIS 147

QY 121 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 165

Db 148 PPDASAAPLRTTADTRKLFRVYSNLRGKLUYGRACRGD 192

RESULT 58

US-10-612-665-22

; Sequence 22, Application US/10612665

; Publication No. US20040122216A1

; GENERAL INFORMATION:

QY ; APPLICANT: Nielsen, J.

Db ; APPLICANT: Pedersen, J.

QY ; APPLICANT: Gerwien, J.

Db ; APPLICANT: Bay, K.

QY ; APPLICANT: Pedersen, L.

Db ; APPLICANT: Cerami, A.

QY ; APPLICANT: Cerami, C.

Db ; TITLE OF INVENTION: RECOMBINANT TISSUE PROTECTIVE CYTOKINES AND ENCODING NUCLEIC ACIDS THEREOF FOR PROTECTION, RESTORATION, AND ENHANCEMENT OF TISSUES AND ORGANS

QY ; TITLE OF INVENTION: RECOMBINANT TISSUE PROTECTIVE CYTOKINES AND ENCODING NUCLEIC ACIDS THEREOF FOR PROTECTION, RESTORATION, AND ENHANCEMENT OF TISSUES AND ORGANS

Db ; FILE REFERENCE: 10165-022-999

QY ; CURRENT FILING DATE: 2003-07-01

Db ; PRIOR APPLICATION NUMBER: 60/392,455

QY ; PRIOR FILING DATE: 2002-07-01

Db ; PRIOR APPLICATION NUMBER: 60/393,423

QY ; PRIOR FILING DATE: 2002-07-03

; NUMBER OF SEQ ID NOS: 212
; SOFTWARE: Patentin version 3.2
; SBO ID NO: 112
; LENGTH: 193
; TYPE: PRT
; ORGANISM: Artificial
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: mutein
; US-10-665-112

Query Match 100 %; Score 846; DB 4; Length 193;
Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 60
Db 28 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 87

Qy 61 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 120
Db 88 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 147

Qy 121 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 165
Db 148 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 192

RESULT 60
US-10-676-694-10
; Sequence 10, Application US/10676694
; Publication No. US20040214236A1

GENERAL INFORMATION:
; APPLICANT: Nielsen, M.
; APPLICANT: Gerwien, J.
; APPLICANT: Pedersen, L.
; APPLICANT: Leist, M.
; APPLICANT: Sager, T.
; APPLICANT: Brines, M.
; APPLICANT: Cerami, A.
; APPLICANT: Ghezzi, P.
; APPLICANT: Fiordaliso, F.
; APPLICANT: Fratelli, M.
; APPLICANT: Gido, G.

TITLE OF INVENTION: TISSUE PROTECTIVE CYTOKINE RECEPTOR COMPLEX AND ASSAYS FOR IDENTIFICATION OF INVENTION: TISSUE PROTECTIVE COMPOUNDS
FILE REFERENCE: 10165-027-999
CURRENT APPLICATION NUMBER: US/10/676,694
CURRENT FILING DATE: 2003-09-30
PRIOR APPLICATION NUMBER: 60/465,891
PRIOR FILING DATE: 2003-04-25
NUMBER OF SEQ ID NOS: 212
SOFTWARE: Patentin version 3.2
SEQ ID NO 22
LENGTH: 193
TYPE: PRT
ORGANISM: Artificial
FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: mutein
; US-10-676-694-22

Query Match 100 %; Score 846; DB 4; Length 193;
Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 60
Db 28 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 87

Qy 61 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 120
Db 88 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 147

Qy 121 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 165
Db 148 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 192

RESULT 61
US-10-676-694-22
; Sequence 22, Application US/10676694
; Publication No. US20040214236A1

GENERAL INFORMATION:
; APPLICANT: Nielsen, M.
; APPLICANT: Gerwien, J.
; APPLICANT: Pedersen, L.
; APPLICANT: Leist, M.
; APPLICANT: Sager, T.
; APPLICANT: Brines, M.
; APPLICANT: Cerami, A.
; APPLICANT: Ghezzi, P.
; APPLICANT: Fiordaliso, F.
; APPLICANT: Fratelli, M.
; APPLICANT: Gido, G.

TITLE OF INVENTION: TISSUE PROTECTIVE CYTOKINE RECEPTOR COMPLEX AND ASSAYS FOR IDENTIFICATION OF INVENTION: TISSUE PROTECTIVE COMPOUNDS
FILE REFERENCE: 10165-027-999
CURRENT APPLICATION NUMBER: US/10/676,694
CURRENT FILING DATE: 2003-09-30
PRIOR APPLICATION NUMBER: 60/465,891
PRIOR FILING DATE: 2003-04-25
NUMBER OF SEQ ID NOS: 212
SOFTWARE: Patentin version 3.2
SEQ ID NO 22
LENGTH: 193
TYPE: PRT
ORGANISM: Artificial
FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: mutein
; US-10-676-694-22

Query Match 100 %; Score 846; DB 4; Length 193;
Best Local Similarity 100.0%; Pred. No. 1. 8e-85; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 60
Db 28 APPRLICDSRVLERYLEAKEARENITGCAEHCSLENITVPTDKNFIYAKRMVEQQA 87

Qy 61 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 120
Db 88 VEWQGALLSBAVRGQALINNSQPWEPLQHVKAVGSLRSLLTIRLAGQAKEAIS 147

Qy 121 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 165
Db 148 PPDAASAPLTTTADTPRKURFVSNFLRGKLUYTGEACRTGD 192

RESULT 62
US-10-676-694-112
; Sequence 112, Application US/10676694
; Publication No. US20040214236A1

GENERAL INFORMATION:
; APPLICANT: Nielsen, M.
; APPLICANT: Gerwien, J.
; APPLICANT: Pedersen, L.
; APPLICANT: Leist, M.
; APPLICANT: Sager, T.
; APPLICANT: Brines, M.
; APPLICANT: Cerami, A.
; APPLICANT: Ghezzi, P.
; APPLICANT: Fiordaliso, F.
; APPLICANT: Fratelli, M.
; APPLICANT: Gido, G.

TITLE OF INVENTION: TISSUE PROTECTIVE CYTOKINE RECEPTOR COMPLEX AND ASSAYS FOR IDENTIFICATION OF INVENTION: TISSUE PROTECTIVE COMPOUNDS
FILE REFERENCE: 10165-027-999
CURRENT APPLICATION NUMBER: US/10/676,694
CURRENT FILING DATE: 2003-09-30

PRIOR APPLICATION NUMBER: 60/465,891 ;
; PRIOR FILING DATE: 2003-04-25 ;
; NUMBER OF SEQ ID NOS: 212 ;
; SOFTWARE: PatentIn version 3.2 ;
; SEQ ID NO 112 ;
; LENGTH: 193 ;
; TYPE: PRT ;
; ORGANISM: Artificial .
; OTHER INFORMATION: Description of Artificial Sequence: mutein
; US-10-676-694-112 ;
; Query Match 100.0%; Score 846; DB 4; Length 193;
; Best Local Similarity 100.0%; Pred. No. 1.8e-85; ;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; SEQ ID NO 1 APPRLICDSRVLERYLLEAKENAINTTGCAEHCSLNENITVPDTKVNFKWAKRNEVGQAA 60
; Db 28 APPRLICDSRVLERYLLEAKENAINTTGCAEHCSLNENITVPDTKVNFKWAKRNEVGQAA 87
; Qy 61 VEWVQGLALLSEAVLRGQALVNSSQPWEPLQLHVDKAVSGRLSLLTIRALGAOKAIS 120
; Db 88 VEWVQGLALLSEAVLRGQALVNSSQPWEPLQLHVDKAVSGRLSLLTIRALGAOKAIS 147
; Qy 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 165
; Db 148 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 192
; RESULT 63
; US-10-759-031-10 ;
; Sequence 10. Application US/10759031
; Publication No. US20050158822A1
; GENERAL INFORMATION:
; APPLICANT: Pecker, Trib
; TITLE OF INVENTION: HIGH LEVEL EXPRESSION OF RECOMBINANT HUMAN ERYTHROPOIETIN
; TITLE OF INVENTION: HAVING
; TITLE OF INVENTION: A MODIFIED 5'-UTR
; FILE REFERENCE: 27179
; CURRENT APPLICATION NUMBER: US/10/759,031
; CURRENT FILING DATE: 2004-01-20
; NUMBER OF SEQ ID NOS: 13
; SOFTWARE: PatentIn version 3.2
; SEQ ID NO 10 ;
; LENGTH: 193
; TYPE: PRT ;
; ORGANISM: Homo sapiens
; US-10-759-031-10
; Query Match 100.0%; Score 845; DB 5; Length 193;
; Best Local Similarity 100.0%; Pred. No. 1.8e-85; ;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; SEQ ID NO 1 APPRLICDSRVLERYLLEAKENAINTTGCAEHCSLNENITVPDTKVNFKWAKRNEVGQAA 60
; Db 28 APPRLICDSRVLERYLLEAKENAINTTGCAEHCSLNENITVPDTKVNFKWAKRNEVGQAA 87
; Qy 61 VEWVQGLALLSEAVLRGQALVNSSQPWEPLQLHVDKAVSGRLSLLTIRALGAOKAIS 120
; Db 88 VEWVQGLALLSEAVLRGQALVNSSQPWEPLQLHVDKAVSGRLSLLTIRALGAOKAIS 147
; Qy 121 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 165
; Db 148 PPDASAAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 192
; RESULT 65
; US-11-021-516-14 ;
; Sequence 14. Application US/11021516
; Publication No. US20050170457A1
; GENERAL INFORMATION:
; APPLICANT: Centocor, Inc.
; APPLICANT: Cunningham, Mark
; APPLICANT: Mills, Julianne
; APPLICANT: Pool, Chandler
; TITLE OF INVENTION: NOVEL RECOMBINANT PROTEINS WITH N-TERMINAL FREE THIOL
; FILE REFERENCE: CEN 5046
; CURRENT APPLICATION NUMBER: US/11/021,516
; CURRENT FILING DATE: 2004-12-23
; PRIOR APPLICATION NUMBER: 60/533617
; PRIOR FILING DATE: 2003-12-31
; NUMBER OF SEQ ID NOS: 20
; SOFTWARE: PatentIn version 3.3
; SEQ ID NO 14
; LENGTH: 193
; TYPE: PRT ;
; ORGANISM: Homo sapiens
; FEATURE:
; NAME/KEY: VARIANT
; LOCATION: (22)..(22)
; OTHER INFORMATION: Q22R
; US-11-021-516-14
; Query Match 100.0%; Score 846; DB 6; Length 193;
; Best Local Similarity 100.0%; Pred. No. 1.8e-85; ;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

1 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 60
 Qy |||||||
 Db ||||||| LENGTH: 209
 28 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 87
 ; FEATURE: TYPE: PRT
 ; ORGANISM: Artificial Sequence
 ; OTHER INFORMATION: Description of Artificial Sequence: Fusion protein
 ; FILE REFERENCE: (BSP) of erythropoietin (EPO) and carboxy terminal
 ; OTHER INFORMATION: peptide (STP) of human thrombopoietin
 Qy |||||||
 Db 61 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 120
 88 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 147
 121 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 165
 148 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 192
 RESULT 66
 US-11-021-516-20
 ; Sequence 20, Application US/11021516
 ; Publication No. US20050170457A1
 ; GENERAL INFORMATION:
 ; APPLICANT: Centocor, Inc.
 ; APPLICANT: Cunningham, Mark
 ; APPLICANT: Mills, Julianne
 ; APPLICANT: Pool, Chandler
 ; TITLE OF INVENTION: NOVEL RECOMBINANT PROTEINS WITH N-TERMINAL FRER THIOL
 ; FILE REFERENCE: CEN 5046
 ; CURRENT APPLICATION NUMBER: US/11/021, 516
 ; CURRENT FILING DATE: 2004-12-23
 ; PRIORITY APPLICATION NUMBER: 60/533617
 ; SEQ ID NO: 20
 ; NUMBER OF SEQ ID NOS: 20
 ; SOFTWARE: Patentin version 3.3
 ; PRIOR FILING DATE: 2003-12-31
 ; LENGTH: 201
 ; TYPE: PRT
 ; ORGANISM: Homo sapiens
 US-11-021-516-20
 Query Match Score 846; DB 6; Length 201;
 Best Local Similarity 100.0%; Pred. No. 1.9e-85; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mi-matches 0;
 Oy 1 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 60
 Db 28 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 87
 Qy |||||||
 61 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 120
 88 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 147
 121 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 165
 148 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 192
 RESULT 67
 US-10-230-454-4
 ; Sequence 4, Application US/10230454
 ; Publication No. US20030124115A1
 ; GENERAL INFORMATION:
 ; APPLICANT: DONG-EOK, LEE
 ; APPLICANT: KYUNG-SUK, OH
 ; APPLICANT: BO-SUP, CHUNG
 ; APPLICANT: JI-SOOK, PARK
 ; APPLICANT: KI-WAN, KIM
 ; TITLE OF INVENTION: FUSION PROTEIN HAVING ENHANCED IN VIVO ACTIVITY OF
 ; TITLE OF INVENTION: BRYTHROPOETIN
 ; FILE REFERENCE: 58105 (71970)
 ; CURRENT APPLICATION NUMBER: US/10/230, 454
 ; CURRENT FILING DATE: 2002-08-29
 ; PRIORITY APPLICATION NUMBER: 2001-74975
 ; PRIOR FILING DATE: 2001-11-29
 ; NUMBER OF SEQ ID NOS: 18
 SOFTWARE: Patentin Ver. 2.1
 RESULT 67
 US-10-230-454-4
 ; Sequence 4, Application US/10230454
 ; Publication No. US20030124115A1
 ; GENERAL INFORMATION:
 ; APPLICANT: DONG-EOK, LEE
 ; APPLICANT: KYUNG-SUK, OH
 ; APPLICANT: BO-SUP, CHUNG
 ; APPLICANT: JI-SOOK, PARK
 ; APPLICANT: KI-WAN, KIM
 ; TITLE OF INVENTION: FUSION PROTEIN HAVING ENHANCED IN VIVO ACTIVITY OF
 ; TITLE OF INVENTION: BRYTHROPOETIN
 ; FILE REFERENCE: 58105 (71970)
 ; CURRENT APPLICATION NUMBER: US/10/230, 454
 ; CURRENT FILING DATE: 2002-08-29
 ; PRIORITY APPLICATION NUMBER: 2001-74975
 ; PRIOR FILING DATE: 2001-11-29
 ; NUMBER OF SEQ ID NOS: 18
 SOFTWARE: Patentin Ver. 2.1
 RESULT 68
 US-10-196-183-2
 ; Sequence 2, Application US/10196183
 ; Publication No. US20030113871A1
 ; GENERAL INFORMATION:
 ; APPLICANT: Lee, Dong-eok
 ; APPLICANT: Park, Ji-sook
 ; APPLICANT: Chung, Bo-sup
 ; APPLICANT: Kim, Ki-wan
 ; APPLICANT: Oh, Myung-suk
 ; TITLE OF INVENTION: Fusion protein having an enhanced in vivo erythropoietin activi
 ; FILE REFERENCE: 401729/YPBR
 ; CURRENT APPLICATION NUMBER: US/10/196, 183
 ; CURRENT FILING DATE: 2002-07-17
 ; PRIORITY APPLICATION NUMBER: KR 10-2001-75994
 ; PRIOR FILING DATE: 2001-12-03
 ; NUMBER OF SEQ ID NOS: 10
 ; SOFTWARE: Patentin version 3.1
 ; SEQ ID NO: 2
 ; LENGTH: 220
 ; TYPE: PRT
 ; ORGANISM: Artificial Sequence
 ; FEATURE:
 ; OTHER INFORMATION: Fusion protein (EATP) of erythropoietin (EPO) and a variant of
 ; OTHER INFORMATION: arboxy terminal peptide (ATP) of human chorionic gonadotropin (H
 ; OTHER INFORMATION: G) beta subunit
 US-10-196-183-2
 Query Match Score 846; DB 4; Length 220;
 Best Local Similarity 100.0%; Pred. No. 2.2e-85; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mi-matches 0;
 Oy 1 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 60
 Db 28 APPRLICDSRVLERYLLEAKAENTITGCAEHCSLENITVPTDKYFVAKRMEVGQA 87
 Qy |||||||
 61 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 120
 88 VEWQGLALLSEAVLRGQALLVNSSQWPEPLQHDKAVSGLRSITLRLGAQEAIIS 147
 121 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 165
 148 PPDAASAPLRTTADTPRKFLFRYVSNFLRGKLUYTGEACRTGD 192
 RESULT 69

US-10-230-454-3
; Sequence 3, Application US/10230454
; Publication No. US20030124115A1
; GENERAL INFORMATION:
; APPLICANT: DONG-EOK, LEE
; APPLICANT: MYUNG-SIK, OH
; APPLICANT: BO-SUP, CHUNG
; APPLICANT: JI-SOOK, PARK
; APPLICANT: KI-WAN, KIM
; TITLE OF INVENTION: FUSION PROTEIN HAVING ENHANCED IN VIVO ACTIVITY OF FILE REFERENCE: 581.05. (71970)
; CURRENT APPLICATION NUMBER: US/10/230,454
; CURRENT FILING DATE: 2002-08-29
; PRIORITY FILING DATE: 2001-11-29
; NUMBER OF SEQ ID NOS: 18
; SOFTWARE: Patentin Ver. 2.1
; SEQ ID NO 3
; LENGTH: 370
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: Description of Artificial Sequence: Fusion protein
; OTHER INFORMATION: (ELTTP) of erythropoietin (EPO) and carboxy terminal
; OTHER INFORMATION: Peptide (LTP) of human thrombopoietin
; US-10-230-454-3

Query Match 100.0%; Score 846; DB 4; Length 370;
Best Local Similarity 100.0%; Pred. No. 4.5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFTAWKMEVGQA 60
Db 28 APPRLICDSVRLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFTAWKMEVGQA 87

QY 61 VEWMOGLALLSEAVLRQALLVNSQWPBQLQAVDGVSGRSLSITLRLGAQKEAIS 120
Db 88 VEWMOGLALLSEAVLRQALLVNSQWPBQLQAVDGVSGRSLSITLRLGAQKEAIS 147

QY 121 PPDAASAAPRITTADETRKLFRVYSNPLRGKLUYGEACRGD 165
Db 148 PPDAASAAPRITTADETRKLFRVYSNPLRGKLUYGEACRGD 192

RESULT 70
US-11-026-998-14
; Sequence 14, Application US/11026998
; Publication No. US20050192211A1
; GENERAL INFORMATION:
; APPLICANT: Gillies, Stephen D.
; APPLICANT: Lauder, Scott
; TITLE OF INVENTION: FC-ERYTHROPOETIN FUSION PROTEIN WITH IMPROVED PHARMACOKINETICS FILE REFERENCE: LEX-027

CURRENT APPLICATION NUMBER: US/11/026,998
CURRENT FILING DATE: 2004-12-30
PRIORITY FILING DATE: 2003-12-31
NUMBER OF SEQ ID NOS: 24
SOFTWARE: Patentin version 3.3
SEQ ID NO 14
LENGTH: 397
TYPE: PRT
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: An amino acid sequence of Fc-EPO containing FN>AQ mutations.
US-11-027-309A-14

Query Match 100.0%; Score 846; DB 6; Length 397;
Best Local Similarity 100.0%; Pred. No. 4.5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSVRLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFTAWKMEVGQA 60
Db 232 APPRLICDSVRLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFTAWKMEVGQA 291

QY 61 VEWMOGLALLSEAVLRQALLVNSQWPBQLQAVDGVSGRSLSITLRLGAQKEAIS 120
Db 292 VEWMOGLALLSEAVLRQALLVNSQWPBQLQAVDGVSGRSLSITLRLGAQKEAIS 351

QY 121 PPDAASAAPRITTADETRKLFRVYSNPLRGKLUYGEACRGD 165
Db 352 PPDAASAAPRITTADETRKLFRVYSNPLRGKLUYGEACRGD 396

RESULT 72
US-10-435-608-10
; Sequence 10, Application US/10435608
; Publication No. US20030235536A1
; GENERAL INFORMATION:
; APPLICANT: Blumberg, Richard S.
; APPLICANT: Lencer, Wayne I.
; APPLICANT: Sinister, Neil E.
; APPLICANT: Biondi, Alan J.
; TITLE OF INVENTION: CENTRAL AIRWAY ADMINISTRATION FOR SYSTEMIC DELIVERY OF THERAPEUT FILE REFERENCE: S01383_70010_US
; CURRENT APPLICATION NUMBER: US/10/435,608
; CURRENT FILING DATE: 2003-05-09
; PRIORITY FILING DATE: 2002-07-03
; NUMBER OF SEQ ID NOS: 27

Query Match 100.0%; Score 846; DB 6; Length 397;
Best Local Similarity 100.0%; Pred. No. 4.5e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
OTHER INFORMATION: An amino acid sequence of Fc-EPO containing FN>AQ mutations.
US-11-026-998-14

; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 10
; LENGTH: 428
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-435-608-10

Query Match 100.0%; Score 846; DB 4; Length 428;
Best Local Similarity 100.0%; Pred. No. 5.5e-05; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 60
Db 28 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 87

Qy 61 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 120
Db 88 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 147

Qy 121 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 165
Db 148 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 192

RESULT 73
US-10-622-108-10
; Sequence 10, Application US/10622108
; Publication No. US20040063912A1
; GENERAL INFORMATION:
; APPLICANT: Blumberg, Richard S.
; APPLICANT: Lencer, Wayne I.
; APPLICANT: Simister, Neil E.
; APPLICANT: Bitonti, Alan J.
; TITLE OF INVENTION: CENTRAL AIRWAY ADMINISTRATION FOR SYSTEMIC DELIVERY OF THERAPEUTIC FILE REFERENCE: S01383_7001.US
; CURRENT APPLICATION NUMBER: US/10/622.108
; CURRENT FILING DATE: 2003-07-17
; PRIOR APPLICATION NUMBER: US 10/435, 608
; PRIOR FILING DATE: 2003-05-09
; PRIOR APPLICATION NUMBER: PCT/US02/21355
; PRIOR FILING DATE: 2002-07-03
; PRIOR APPLICATION NUMBER: US 60/364, 482
; PRIOR FILING DATE: 2002-03-15
; NUMBER OF SEQ ID NOS: 40
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 10
; LENGTH: 428
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-622-108-10

Query Match 100.0%; Score 846; DB 4; Length 428;
Best Local Similarity 100.0%; Pred. No. 5.5e-05; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 60
Db 28 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 87

Qy 61 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 120
Db 88 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 147

Qy 121 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 165
Db 148 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 192

RESULT 74
US-10-841-250-24
; Sequence 22, Application US/09932812
; Publication No. US20030082749A1
; GENERAL INFORMATION:
; APPLICANT: Sun, Iee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological activity FILE REFERENCE: US20030082749A1
; CURRENT FILING DATE: 2001-10-30
; NUMBER OF SEQ ID NOS: 22
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 22
; LENGTH: 435
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURES:
; OTHER INFORMATION: HubPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure 2)
; US-09-932-812-22

Query Match 100.0%; Score 846; DB 3; Length 435;
Best Local Similarity 100.0%; Pred. No. 5.6e-05; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 60
Db 28 APPRLICDSRVLERYLLEAKRKENITGCAHCISLENNTIPDTKVNFYAWKRMVQQA 87

Qy 61 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 120
Db 88 VEWQGLALLSBAVLRGOALLVNSSQEPWPLQLHVDKAVSGRSLSLTLLRAIGAQEAI 147

Qy 121 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 165
Db 148 PPDAASAPLRTTADTPRKLFRVYNSFLRGKLYTGAEACTGD 192

RESULT 74
US-10-841-250-24
; Sequence 24, Application US/10841250
; Publication No. US20050032174A1
; GENERAL INFORMATION:

RESULT 76 ; FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 120 ; OTHER INFORMATION: HUEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-10-761-593A-22)

Db ; Sequence 22, Application US/10761593A

Publication No. US20040175824A1

GENERAL INFORMATION:

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with high biological

FILE REFERENCE: 02SUM2001-A

CURRENT APPLICATION NUMBER: US/10/761,593A

PRIOR APPLICATION NUMBER: 09/932812

PRIOR FILING DATE: 2004-01-21

NUMBER OF SEQ ID NOS: 28

SOFTWARE: PatentIn version 3.2

SEQ ID NO: 22

LENGTH: 435

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 147

OTHER INFORMATION: HuEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-10-761-593A-22)

Query Match 100.0% Score 846; DB 4; Length 435;

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

RESULT 77 ; FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 120 ; OTHER INFORMATION: HUEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-11-016-518A-22)

Db ; Sequence 22, Application US/11016518A

Publication No. US20050142642A1

GENERAL INFORMATION:

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological

FILE REFERENCE: 02SUM2001D2

CURRENT APPLICATION NUMBER: US/11/017,185

PRIOR APPLICATION NUMBER: US 09/932,812

PRIOR FILING DATE: 2004-12-17

NUMBER OF SEQ ID NOS: 28

SOFTWARE: PatentIn version 3.1

SEQ ID NO: 22

LENGTH: 435

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 147

OTHER INFORMATION: HuEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-11-016-518A-22)

Query Match 100.0% Score 846; DB 6; Length 435;

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

RESULT 78 ; FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 120 ; OTHER INFORMATION: HUEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-11-017-185-22)

Db ; Sequence 22, Application US/11017185

Publication No. US20050142642A1

GENERAL INFORMATION:

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological

FILE REFERENCE: 02SUM2001D2

CURRENT APPLICATION NUMBER: US/11/017,185

PRIOR APPLICATION NUMBER: US 09/932,812

PRIOR FILING DATE: 2004-12-17

NUMBER OF SEQ ID NOS: 28

SOFTWARE: PatentIn version 3.1

SEQ ID NO: 22

LENGTH: 435

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 147

OTHER INFORMATION: HuEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-11-017-185-22)

Query Match 100.0% Score 846; DB 6; Length 435;

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

Best local similarity 100.0%; Pred. No. 5.6e-85; Mismatches 0; Indels 0; Gaps 0; Matches 165; Conservative

RESULT 79 ; FEATURE: VEWQGLALISEAVRLRQALLVNSQPWEPLQLANDKAVSGRLSITLRLAAGQKEAIS 120 ; OTHER INFORMATION: HUEPO-L-vFc gamma1 with a 27-amino acid leader peptide (Figure US-09-932-812-18)

Db ; Sequence 18, Application US/09932812

Publication No. US20030082749A1

GENERAL INFORMATION:

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased

FILE REFERENCE: 02SUM2004D1

CURRENT APPLICATION NUMBER: US/11/016,518A

PRIOR APPLICATION NUMBER: US 09/932,812

PRIOR FILING DATE: 2004-12-17

NUMBER OF SEQ ID NOS: 28

SOFTWARE: PatentIn version 3.2

SEQ ID NO: 22

LENGTH: 435

TYPE: PRT

ORGANISM: Artificial Sequence

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological activities

FILE REFERENCE: 02SUN2001

CURRENT APPLICATION NUMBER: US/09/932,812

CURRENT FILING DATE: 2001-10-30

NUMBER OF SEQ ID NOS: 22

SEQ ID NO 18

LENGTH: 436

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: OTHER INFORMATION: HuEPO-L-vFc gamma2 with a 27-amino acid leader peptide (Figure 2)

OTHER INFORMATION: A)

US-09-932-812-18

Query Match 100.0%; Score 846; DB 3; Length 436;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 60

Db 28 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 87

QY 61 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 120

Db 88 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 147

QY 121 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 165

Db 148 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 192

RESULT 80

US-10-761-593A-18
Sequence 18 Application US/10761593A

GENERAL INFORMATION:
Publication No. US200401758241

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with high biological

TITLE OF INVENTION: activities

FILE REFERENCE: 02SUN2001-A

CURRENT APPLICATION NUMBER: US/10/761,593A

PRIOR APPLICATION NUMBER: 09/932812

PRIOR FILING DATE: 2001-08-17

NUMBER OF SEQ ID NOS: 28

SEQ ID NO 18

LENGTH: 436

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: OTHER INFORMATION: HuEPO-L-vFc gamma2 with a 27-amino acid leader peptide (Figure

US-10-761-593A-18

Query Match 100.0%; Score 846; DB 6; Length 436;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 60

Db 28 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 87

QY 61 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 120

Db 88 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 147

QY 121 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 165

Db 148 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 192

RESULT 82

US-11-017-185-18
Sequence 18 Application US/11017185

GENERAL INFORMATION:
Publication No. US20050142642A1

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological

TITLE OF INVENTION: activities

FILE REFERENCE: 02SUN2001D2

CURRENT APPLICATION NUMBER: US/11/017,185

PRIOR APPLICATION NUMBER: US 09/932,812

PRIOR FILING DATE: 2001-08-17

NUMBER OF SEQ ID NOS: 28

SEQ ID NO 18

LENGTH: 436

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: OTHER INFORMATION: HuEPO-L-vFc gamma2 with a 27-amino acid leader peptide (Figure

US-11-017-185-18

Query Match 100.0%; Score 846; DB 6; Length 436;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 60

Db 28 APPRLICDSRVRVLRVYLLEAKAENITVGCAEHCSLNENITVPDTKVNPFYANKRMEVGQAA 87

QY 61 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 120

Db 88 VEWMOGLLISSEAVLRLGQALLVNSQPWPQLQHVDKAVSGLSLTTLRALGAQKEAIS 147

QY 121 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 165

Db 148 PPDASAPLRTTADTRKLFRVSNFLRGKLUYGEACRTGD 192

RESULT 81

US-11-016-18A-18

Sequence 18 Application US/11016518A

Publication No. US20050124045A1

GENERAL INFORMATION:

APPLICANT: Sun, Lee-Hwei K

APPLICANT: Sun, Bill N

APPLICANT: Sun, Cecily R

TITLE OF INVENTION: Fc fusion proteins of human erythropoietin with increased biological

TITLE OF INVENTION: activities

FILE REFERENCE: 02SUN2004D1

CURRENT APPLICATION NUMBER: US/11/016,518A

PRIOR APPLICATION NUMBER: US 09/932,812

PRIOR FILING DATE: 2004-12-17

NUMBER OF SEQ ID NOS: 28

SEQ ID NO 18

LENGTH: 436

TYPE: PRT

ORGANISM: Artificial Sequence

FEATURE: OTHER INFORMATION: HuEPO-L-vFc gamma2 with a 27-amino acid leader peptide (Figure

US-11-017-185-18

Query Match 100.0%; Score 846; DB 6; Length 436;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 60
Db 28 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 87

QY 61 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 120
Db 88 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 147

QY 121 PPDASAAPLRTTADTFRKLFVRYSNFLRGKLUYTGEACRTGD 165
Db 148 PPDASAAPLRTTADTFRKLFVRYSNFLRGKLUYTGEACRTGD 192

RESULT 83
US-09-932-812-20
; Sequence 20, Application US/0932812
; Publication No. US20030082749A1
; GENERAL INFORMATION:
; APPLICANT: Sun, Lee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: FC fusion proteins of human erythropoietin with increased biological activity

FILE REFERENCE: 02SUN2001
CURRENT APPLICATION NUMBER: US/09/932, 812
CURRENT FILING DATE: 2001-10-30
NUMBER OF SEQ ID NOS: 22
SOFTWARE: PatentIn version 3.1
SEQ ID NO 20
LENGTH: 437
TYPE: PRT
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: HUEPO-L-vFC gamma4 with a 27-amino acid leader peptide (Figure 2B)

Query Match 100.0%; Score 846; DB 3; Length 437;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 60
Db 28 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 87

QY 61 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 120
Db 88 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 147

RESULT 85
US-11-016-518A-20
; Sequence 20, Application US/11016518A
; Publication No. US20050124045A1
; GENERAL INFORMATION:
; APPLICANT: Sun, Lee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: FC fusion proteins of human erythropoietin with increased biological activity

FILE REFERENCE: 02SUN2004DI
CURRENT APPLICATION NUMBER: US/11/016, 518A
CURRENT FILING DATE: 2004-12-17
PRIORITY FILING DATE: 2001-08-17
NUMBER OF SEQ ID NOS: 28
SOFTWARE: PatentIn version 3.2
SEQ ID NO 20
LENGTH: 437
TYPE: PRT
ORGANISM: Artificial Sequence
FEATURE:
OTHER INFORMATION: HUEPO-L-vFC gamma4 with a 27-amino acid leader peptide (Figure 2B)

Query Match 100.0%; Score 846; DB 6; Length 437;
Best Local Similarity 100.0%; Pred. No. 5.6e-85;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 60
Db 28 APPRICDSRVLYLEAKEAENITGCAEHCSLENITVPDKVNFKWAKMENVGQA 87

QY 61 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 120
Db 88 VEWQGLALLSEAVLRGQALLVNSQWPMLQHDKAVSGLRSLTTLRAGQKEAIS 147

RESULT 86
US-11-017-185-20
; Sequence 20, Application US/11017185
; Publication No. US20050142642A1
; GENERAL INFORMATION:

CURRENT APPLICATION NUMBER: US/10/761, 593A
CURRENT FILING DATE: 2004-01-21
PRIOR APPLICATION NUMBER: 09/932812
NUMBER OF SEQ ID NOS: 28
SOFTWARE: PatentIn version 3.2
SEQ ID NO 20

APPLICANT: Sun, Lee-Hwei K ; SOFTWARE: Patentin Ver. 2.0
; APPLICANT: Sun, Bill N ; SEQ ID NO 1521
; APPLICANT: Sun, Cecily R ; LENGTH: 768
; TITLE OF INVENTION: fc fusion proteins of human erythropoietin with increased biologi
; TITLE OF INVENTION: activities
; FILE REFERENCE: 02SUN2001D2
; CURRENT APPLICATION NUMBER: US/11/017,185
; CURRENT FILING DATE: 2004-12-17
; PRIOR APPLICATION NUMBER: US 09/932,812
; NUMBER OF SEQ ID NOS: 28
; SEQ ID NO 20
; LENGTH: 437
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: HubPO-L-vPC gamma4 with a 27-amino acid leader peptide (Figure 2E)
; OTHER INFORMATION:)
; OTHER INFORMATION: US-11-017-185-20

Query Match 100%; Score 846; DB 5; Length 437;
Best Local Similarity 100.0%; Pred. No. 5. 6e-05; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFIWAKRMEVGQQA 60
Db 28 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFIWAKRMEVGQQA 87

QY 61 VEWMOGLALLSBAVLRGQALVNSSQPEPLQLHVDKAVSGRSLTILRALGAQKRAIS 120
Db 88 VEWMOGLALLSBAVLRGQALVNSSQPEPLQLHVDKAVSGRSLTILRALGAQKRAIS 147

QY 121 PPDAASAPRLTTADPFKLFRVYSNPLRGKLUYTGEACRTGD 165
Db 148 PPDAASAPRLTTADPFKLFRVYSNPLRGKLUYTGEACRTGD 192

RESULT 87

US-10-775-204-1521

; Sequence 1521, Application US/10775204
; Publication No. US200501866641

; GENERAL INFORMATION:

; APPLICANT: Rosen, Craig A.
; APPLICANT: Baseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: P564
; CURRENT APPLICATION NUMBER: US/10/775,204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341,811
; PRIOR FILING DATE: 2002-12-21
; PRIOR APPLICATION NUMBER: 60/360,000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378,950
; PRIOR FILING DATE: 2002-05-10
; PRIOR APPLICATION NUMBER: 60/398,008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411,355
; PRIOR FILING DATE: 2002-09-18
; PRIOR APPLICATION NUMBER: 60/414,984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/417,611
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/420,246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423,623
; PRIOR FILING DATE: 2002-11-05
; PRIOR APPLICATION NUMBER: 60/351,360
; PRIOR FILING DATE: 2002-01-28
; remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: Patentin ver. 2.0
; SEQ ID NO 1522
; LENGTH: 768
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-1522

Query Match 100%; Score 846; DB 5; Length 768;
Best Local Similarity 100.0%; Pred. No. 1. 2e-84; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFIWAKRMEVGQQA 60
Db 604 APPRLICDSRVLERYLLEAKEAENITTGCAEHCSLNENITVPDTKVNFIWAKRMEVGQQA 663

Qy 61 VEWVOGLALISEAVRGOALLVNSQPEPLQLHVDAVSGLSLTILRAIGAQEAI S 120
Db 664 VEWVOGLALISEAVRGOALLVNSQPEPLQLHVDAVSGLSLTILRAIGAQEAI S 723

Qy 121 PPDASAAPLRTTADTPKLFRYTSNFLAGKLYTGACRTGD 165
Db 724 PPDASAAPLRTTADTPKLFRYTSNFLAGKLYTGACRTGD 768

RESULT 89
US-10-775-204-1523
; Sequence 1523, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Haseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: PF564
; CURRENT APPLICATION NUMBER: US/10/775, 204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341, 811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/3360, 000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378, 950
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341, 811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/360, 000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378, 950
; PRIOR FILING DATE: 2002-03-10
; PRIOR APPLICATION NUMBER: 60/398, 008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411, 355
; PRIOR FILING DATE: 2002-03-18
; PRIOR APPLICATION NUMBER: 60/414, 984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/417, 611
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/420, 246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423, 623
; PRIOR FILING DATE: 2002-11-05
; PRIOR APPLICATION NUMBER: 60/351, 360
; PRIOR FILING DATE: 2002-01-28
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 1660
; LENGTH: 768
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-1523
; ORGANISM: Homo sapiens
Query Match 100.0%; Score 846; DB 5; Length 768;
Best Local Similarity 100.0%; Pred. No. 1.2e-84; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Query Match 100.0%; Score 846; DB 5; Length 768;
; Best Local Similarity 100.0%; Pred. No. 1.2e-84; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 APRLICDSRVLYERVLKEAKENITTCGAECISLNENITVPDTKVNFWKRMVGQA 60
Db 604 APRLICDSRVLYERVLKEAKENITTCGAECISLNENITVPDTKVNFWKRMVGQA 663

Qy 61 VEWVOGLALISEAVRGOALLVNSQPEPLQLHVDAVSGLSLTILRAIGAQEAI S 120
Db 664 VEWVOGLALISEAVRGOALLVNSQPEPLQLHVDAVSGLSLTILRAIGAQEAI S 723

Qy 121 PPDASAAPLRTTADTPKLFRYTSNFLAGKLYTGACRTGD 165
Db 724 PPDASAAPLRTTADTPKLFRYTSNFLAGKLYTGACRTGD 768

RESULT 91
US-10-775-204-1661
; Sequence 1661, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Haseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: PF564
; CURRENT APPLICATION NUMBER: US/10/775, 204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341, 811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/360, 000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378, 950
; PRIOR FILING DATE: 2002-03-10
; PRIOR APPLICATION NUMBER: 60/398, 008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411, 355
; PRIOR FILING DATE: 2002-09-18
; PRIOR APPLICATION NUMBER: 60/414, 984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/417, 611
; PRIOR FILING DATE: 2002-10-11
; PRIOR APPLICATION NUMBER: 60/420, 246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423, 623
; PRIOR FILING DATE: 2002-11-05
; PRIOR APPLICATION NUMBER: 60/351, 360
; PRIOR FILING DATE: 2002-01-28
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 1660
; LENGTH: 768
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-1660

RESULT 90
US-10-775-204-1660
; Sequence 1660, Application US/10775204
; Publication No. US20050186664A1

PRIOR FILING DATE: 2002-05-10 ;
; PRIOR APPLICATION NUMBER: 60/398,008
; PRIOR FILING DATE: 2002-07-24 ;
; PRIOR APPLICATION NUMBER: 60/411,355
; PRIOR FILING DATE: 2002-09-18 ;
; PRIOR APPLICATION NUMBER: 60/414,984
; PRIOR FILING DATE: 2002-10-02 ;
; PRIOR APPLICATION NUMBER: 60/417,611
; PRIOR FILING DATE: 2002-10-11 ;
; PRIOR APPLICATION NUMBER: 60/420,246
; PRIOR FILING DATE: 2002-10-23 ;
; SEQ ID NO: 1662
; LENGTH: 768
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-1661

Query Match 100.0%; Score 846; DB 5; Length 768;
Best Local Similarity 100.0%; Pred. No. 1.2e-84; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITTGCAECSLENNTVDPDTKVNFYAWKRMEMVGQA 60
Db 604 APPRLICDSRVLERYLLEAKEAENITTGCAECSLENNTVDPDTKVNFYAWKRMEMVGQA 663
; Sequence 1662, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Haseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: P2564
; CURRENT APPLICATION NUMBER: US/10/775,204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341,811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/360,000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378,950
; PRIOR FILING DATE: 2002-05-10
; PRIOR APPLICATION NUMBER: 60/398,008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411,355
; PRIOR FILING DATE: 2002-09-18
; PRIOR APPLICATION NUMBER: 60/414,984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/420,246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423,623
; PRIOR FILING DATE: 2002-11-05
; SEQ ID NO: 387
; LENGTH: 769
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-387

Query Match 100.0%; Score 846; DB 5; Length 769;
Best Local Similarity 100.0%; Pred. No. 1.2e-84; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

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Db 604 APPRLICDSRVLERYLLEAKEAENITTGCAECSLENNTVDPDTKVNFYAWKRMEMVGQA 663
; Sequence 1662, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Haseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: P2564
; CURRENT APPLICATION NUMBER: US/10/775,204
; CURRENT FILING DATE: 2004-02-11
; PRIOR APPLICATION NUMBER: 60/341,811
; PRIOR FILING DATE: 2001-12-21
; PRIOR APPLICATION NUMBER: 60/360,000
; PRIOR FILING DATE: 2002-02-28
; PRIOR APPLICATION NUMBER: 60/378,950
; PRIOR FILING DATE: 2002-05-10
; PRIOR APPLICATION NUMBER: 60/398,008
; PRIOR FILING DATE: 2002-07-24
; PRIOR APPLICATION NUMBER: 60/411,355
; PRIOR FILING DATE: 2002-09-18
; PRIOR APPLICATION NUMBER: 60/414,984
; PRIOR FILING DATE: 2002-10-02
; PRIOR APPLICATION NUMBER: 60/420,246
; PRIOR FILING DATE: 2002-10-23
; PRIOR APPLICATION NUMBER: 60/423,623
; PRIOR FILING DATE: 2002-11-05
; PRIOR APPLICATION NUMBER: 60/351,360
; PRIOR FILING DATE: 2002-01-28
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO: 387
; LENGTH: 769
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-387

PRIOR APPLICATION NUMBER: 60/360,000 ;
 PRIOR FILING DATE: 2002-02-28 ;
 PRIOR APPLICATION NUMBER: 60/378,950 ;
 PRIOR FILING DATE: 2002-05-10 ;
 PRIOR APPLICATION NUMBER: 60/398,008 ;
 PRIOR FILING DATE: 2002-07-24 ;
 PRIOR APPLICATION NUMBER: 60/411,355 ;
 PRIOR FILING DATE: 2002-09-18 ;
 PRIOR APPLICATION NUMBER: 60/414,984 ;
 PRIOR FILING DATE: 2002-10-02 ;
 PRIOR APPLICATION NUMBER: 60/417,611 ;
 PRIOR FILING DATE: 2002-10-11 ;
 PRIOR APPLICATION NUMBER: 60/420,246 ;
 PRIOR FILING DATE: 2002-10-23 ;
 PRIOR APPLICATION NUMBER: 60/423,623 ;
 PRIOR FILING DATE: 2002-11-05 ;
 PRIOR APPLICATION NUMBER: 60/351,360 ;
 PRIOR FILING DATE: 2002-01-28 ;
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 PRIOR FILING DATE: 2002-01-28 ;
 Remaining Prior Application data removed - See File Wrapper or PALM.
 NUMBER OF SEQ ID NOS: 2222 ;
 SOFTWARE: PatentIn Ver. 2.0 ;
 SEQ ID NO: 375 ;
 LENGTH: 777 ;
 TYPE: PRT ;
 ORGANISM: Homo sapiens ;
 US-10-775-204-374 ;
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 Db 88 VEWMOGLALLSRAVLRGQALLNSSQWPFLQHVDKAVSGRLSLTTLRAGAQEAKIS 147 ;
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 Db 148 PPDASASAPLTTADTFRKLFPRVYSNPLRGKLUYGEACRTGD 192 ;
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 RESULT 97 ;
 US-10-775-204-375 ;
 ; Sequence 375, Application US/10775204 ;
 ; Publication No. US200501866641 ;
 ; GENERAL INFORMATION:
 ; APPLICANT: Rosen, Craig A. ;
 ; ATTORNEY: Haseltine, William A. ;
 ; ATTORNEY: Balance, David J. ;
 ; ATTORNEY: Turner, Andrew J. ;
 ; TITLE OF INVENTION: Albumin Fusion Proteins ;
 ; FILE REFERENCE: P564 ;
 ; CURRENT APPLICATION NUMBER: US/10/775,204 ;
 ; CURRENT FILING DATE: 2004-02-11 ;
 ; PRIOR APPLICATION NUMBER: 60/341,811 ;
 ; PRIOR FILING DATE: 2001-12-21 ;
 ; PRIOR APPLICATION NUMBER: 60/360,000 ;
 ; PRIOR FILING DATE: 2002-02-28 ;
 ; PRIOR APPLICATION NUMBER: 60/378,950 ;
 ; PRIOR FILING DATE: 2002-05-10 ;
 ; PRIOR APPLICATION NUMBER: 60/398,008 ;
 ; PRIOR FILING DATE: 2002-07-24 ;
 ; PRIOR APPLICATION NUMBER: 60/411,355 ;
 ; PRIOR FILING DATE: 2002-09-18 ;
 ; PRIOR APPLICATION NUMBER: 60/414,984 ;
 ; PRIOR FILING DATE: 2002-10-02 ;
 ; PRIOR APPLICATION NUMBER: 60/420,246 ;
 ; PRIOR FILING DATE: 2002-10-23 ;
 ; PRIOR APPLICATION NUMBER: 60/423,623 ;
 ; PRIOR FILING DATE: 2002-11-05 ;
 ; PRIOR APPLICATION NUMBER: 60/351,360 ;
 ; PRIOR FILING DATE: 2002-01-28 ;
 ; Remaining Prior Application data removed - See File Wrapper or PALM.
 NUMBER OF SEQ ID NOS: 2222 ;
 SOFTWARE: PatentIn Ver. 2.0 ;
 SEQ ID NO: 377 ;
 LENGTH: 777 ;
 TYPE: PRT ;
 ORGANISM: Homo sapiens ;
 US-10-775-204-377 ;
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CURRENT FILING DATE: 2004-02-11
; PRIORITY APPLICATION NUMBER: 60/341,811
; PRIORITY FILING DATE: 2001-12-21
; PRIORITY APPLICATION NUMBER: 60/360,000
; PRIORITY FILING DATE: 2002-02-28
; PRIORITY APPLICATION NUMBER: 60/378,950
; PRIORITY FILING DATE: 2002-05-10
; PRIORITY APPLICATION NUMBER: 60/398,008
; PRIORITY FILING DATE: 2002-07-24
; PRIORITY APPLICATION NUMBER: 60/411,355
; PRIORITY FILING DATE: 2002-09-18
; PRIORITY APPLICATION NUMBER: 60/414,984
; PRIORITY FILING DATE: 2002-10-02
; PRIORITY APPLICATION NUMBER: 60/417,611
; PRIORITY FILING DATE: 2002-10-11
; PRIORITY APPLICATION NUMBER: 60/420,246
; PRIORITY FILING DATE: 2002-10-23
; PRIORITY APPLICATION NUMBER: 60/423,623
; PRIORITY FILING DATE: 2002-11-05
; PRIORITY APPLICATION NUMBER: 60/351,360
; PRIORITY FILING DATE: 2002-01-28
; Remaining Prior Application data removed - See File Wrapper or PALM.
; NUMBER OF SEQ ID NOS: 2222
; SOFTWARE: Patentin Ver. 2.0
; SEQ ID NO 409
; LENGTH: 951
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-775-204-409

Query Match 100.0%; Score 846; DB 5; Length 951;
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Qy 121 PPDAASAPLRITTADEPRKLFPRVYNSFLRGKLYTGAEACTGD 165
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RESULT 102

US-10-775-204-401
; Sequence 401, Application US/10775204
; Publication No. US20050186664A1
; GENERAL INFORMATION:
; APPLICANT: Rosen, Craig A.
; APPLICANT: Haseltine, William A.
; APPLICANT: Balance, David J.
; APPLICANT: Turner, Andrew J.
; TITLE OF INVENTION: Albumin Fusion Proteins
; FILE REFERENCE: PP564
; CURRENT APPLICATION NUMBER: US/10/775,204
; CURRENT FILING DATE: 2004-02-11
; PRIORITY APPLICATION NUMBER: 60/341,811
; PRIORITY FILING DATE: 2001-12-21
; PRIORITY APPLICATION NUMBER: 60/360,000
; PRIORITY FILING DATE: 2002-02-28
; PRIORITY APPLICATION NUMBER: 60/378,950
; PRIORITY FILING DATE: 2002-05-10
; PRIORITY APPLICATION NUMBER: 60/398,008
; PRIORITY FILING DATE: 2002-07-24
; PRIORITY APPLICATION NUMBER: 60/411,355
; PRIORITY FILING DATE: 2002-09-18
; PRIORITY APPLICATION NUMBER: 60/414,984
; PRIORITY FILING DATE: 2002-10-02
; PRIORITY APPLICATION NUMBER: 60/417,611

Query Match 100.0%; Score 846; DB 5; Length 954;
Best Local Similarity 100.0%; Pred. No. 1.7e-84; Mismatches 0; Indels 0; Gaps 0;
Matches 165; Conservative 0; MisMatches 0;

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Qy 121 PPDAASAPLRITTADEPRKLFPRVYNSFLRGKLYTGAEACTGD 165
Db 910 PPDAASAPLRITTADEPRKLFPRVYNSFLRGKLYTGAEACTGD 954

Search completed: March 1, 2006, 10:24:34
Job time : 69 secs

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On protein - protein search, using SW model

Run on: February 28, 2006, 15:39:46 ; Search time 18 Seconds
 (without alignments)
 136.466 Million cell updates/sec

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 Gapop 10.0 , Gapext 0.5

Searched: 117670 seqs, 14887254 residues

Total number of hits satisfying chosen parameters: 117670

Minimum DB seq length: 0
 Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
 Maximum Match 100%
 Listing first 45 summaries

Database : Published Applications AA New:
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Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

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1	846	100.0	166	US-10-522-297-1
2	846	100.0	166	Sequence 1, Appli
3	846	100.0	193	Sequence 201, App
4	846	100.0	428	Sequence 4, Appli
5	844	99.8	166	Sequence 24, Appli
6	844	99.8	166	Sequence 959, Appli
7	843	99.6	166	Sequence 967, Appli
8	843	99.6	166	Sequence 952, App
9	843	99.6	166	Sequence 955, App
10	843	99.6	165	Sequence 958, App
11	843	99.6	412	Sequence 966, App
12	843	99.6	444	Sequence 34, Appli
13	842	99.5	166	Sequence 16, Appli
14	842	99.5	166	Sequence 942, App
15	842	99.5	166	Sequence 948, App
16	842	99.5	166	Sequence 951, App
17	842	99.5	166	Sequence 961, App
18	842	99.5	166	Sequence 971, App
19	841	99.4	166	Sequence 941, App
20	841	99.4	166	Sequence 943, App
21	841	99.4	166	Sequence 201, Application US/11176830
22	841	99.4	166	; Publication No. US2006020116A1
23	841	99.4	166	; GENERAL INFORMATION:
24	841	99.4	166	; APPLICANT: Gantier, Rene
25	841	99.4	166	; APPLICANT: Guyon, Thierry

ALIGMENTS

RESULT 1
 US-10-522-297-1
 ; Sequence 1, Application US/10522297
 ; Publication No. US2006020116A1
 ; GENERAL INFORMATION:
 ; APPLICANT: MERCK PATENT GMBH
 ; APPLICANT: BAKER, Matthew
 ; APPLICANT: CARR, Francis J
 ; TITLE OF INVENTION: T-CELL EPITOPES IN ERYTHROPOIETIN
 ; FILE REFERENCE: MBR-137
 ; CURRENT APPLICATION NUMBER: US/10/522,297
 ; CURRENT FILING DATE: 2005-01-24
 ; PRIORITY APPLICATION NUMBER: PCT/EP2003/008725
 ; PRIORITY FILING DATE: 2003-08-07
 ; PRIORITY APPLICATION NUMBER: EP20017914.9
 ; PRIORITY FILING DATE: 2002-08-09
 ; SOFTWARE: FastSEQ for Windows Version 4.0
 ; SEQ ID NO: 61
 ; LENGTH: 166
 ; TYPE: PRT
 ; ORGANISM: Homo sapiens
 ; US-10-522-297-1

RESULT 2
 US-11-176-830-201
 ; Sequence 201, Application US/11176830

Query Match Similarity 100.0%; Pred. No. 1. 2e-84; 0; Mismatches 0; Indels 0; Gaps 0;

Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Organism: Homo sapiens

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Sequence 3 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

Sequence 4 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

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Sequence 19 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

Sequence 20 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

Sequence 21 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

Sequence 22 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

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Sequence 24 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

Sequence 25 APPRLIDSRVLEVLREAKLENITTCGACBICSLNLNENITWDTKVNIFYAWKRMEEVQQA 60

APPLICANT: Drittanti, Lila
 APPLICANT: Vega, Manuel
 TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Nu
 TITLE OF INVENTION: Acid Molecules and Related Applications
 FILE REFERENCE: 17109-012002 (922B)
 CURRENT APPLICATION NUMBER: US11/176, 830
 CURRENT PILING DATE: 2005-07-06
 PRIOR APPLICATION NUMBER: 10/658, 834
 PRIOR FILING DATE: 2003-09-08
 PRIOR APPLICATION NUMBER: 60/457, 135
 PRIOR FILING DATE: 2003-03-21
 PRIOR APPLICATION NUMBER: 60/409, 898
 PRIOR FILING DATE: 2002-09-09
 NUMBER OF SEQ ID NOS: 136
 SOFTWARE: FAST-SEQ for Windows Version 4.0
 SEQ ID NO 201
 LENGTH: 166
 TYPE: PRT
 ORGANISM: Homo sapiens
 PUBLICATION INFORMATION:
 DATABASE ACCESION NUMBER: Genbank AAA52400
 DATABASE ENTRY DATE: 1994-11-08
 US-11-176-830-201

RESULT 4
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 Best Local Similarity 100.0%; Pred. No. 1, 2e-84; Mismatches 0; Indels 0; Gaps 0;
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Qy 61 VEWQGGLALLSEAVRLRGQLVNSSQWPQLHVDKAVSGLRSLTLLRAKGKEAIS 120
 Db 61 VEWQGGLALLSEAVRLRGQLVNSSQWPQLHVDKAVSGLRSLTLLRAKGKEAIS 120

Qy 121 PPDAASAPLRTTADFRKLFRVYSNPLRGKLUYGEACRTGD 165
 Db 121 PPDAASAPLRTTADFRKLFRVYSNPLRGKLUYGEACRTGD 165

RESULT 3
 US-11-144-889A-4
 Sequence 4, Application US/11144889A
 Publication No. US20050272654A1
 GENERAL INFORMATION:
 APPLICANT: HIRSCH, FRANCOIS
 APPLICANT: HARPFNER, ASTRID
 TITLE OF INVENTION: NF-KB ACTIVATION INHIBITORS, AND THEIR PHARMACEUTICAL
 TITLE OF INVENTION: - USES
 FILE REFERENCE: 0508-1046-1
 CURRENT APPLICATION NUMBER: US/11/144, 889A
 CURRENT FILING DATE: 2005-06-06
 PRIOR APPLICATION NUMBER: 09/856, 795
 PRIOR FILING DATE: 2001-05-25
 PRIOR APPLICATION NUMBER: PCT/FR99/02897
 PRIOR FILING DATE: 1999-11-24
 PRIOR APPLICATION NUMBER: FR 99/14858
 PRIOR FILING DATE: 1998-11-25
 NUMBER OF SEQ ID NOS: 4
 SOFTWARE: PatentIn Ver. 3.3
 SEQ ID NO 4
 LENGTH: 193
 TYPE: PRT
 ORGANISM: Homo sapiens
 US-11-144-889A-4

Query Match 100.0%; Score 846; DB 7; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1, 2e-84; Mismatches 0; Indels 0; Gaps 0;
 Matches 165; Conservative 0; Mi mismatches 0; Indels 0; Gaps 0;

Qy 1 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENITVPDTKVNLYAKRMEVGQA 60
 Db 28 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENITVPDTKVNLYAKRMEVGQA 87

Qy 61 VEWQGGLALLSEAVRLRGQLVNSSQWPQLHVDKAVSGLRSLTLLRAKGKEAIS 120
 Db 88 VEWQGGLALLSEAVRLRGQLVNSSQWPQLHVDKAVSGLRSLTLLRAKGKEAIS 147

Qy 121 PPDAASAPLRTTADFRKLFRVYSNPLRGKLUYGEACRTGD 165
 Db 148 PPDAASAPLRTTADFRKLFRVYSNPLRGKLUYGEACRTGD 192

RESULT 5
 US-11-176-830-959
 Sequence 959, Application US/11176830
 Publication No. US20060020116A1
 GENERAL INFORMATION:
 APPLICANT: Gantier, Rene
 APPLICANT: Guyon, Thierry
 APPLICANT: Drittanti, Lila
 APPLICANT: Vega, Manuel
 TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Nu
 TITLE OF INVENTION: Acid Molecules and Related Applications
 FILE REFERENCE: 17109-012002 (922B)
 CURRENT APPLICATION NUMBER: US/11/176, 830
 CURRENT PILING DATE: 2005-07-06

Db 28 APRPLICDSRVLERYLLEAKEAENITTGCAEHCSLENITVPDTKVNLYAKRMEVGQA 87
 Qy 61 VEWQGGLALLSEAVRLRGQLVNSSQWPQLHVDKAVSGLRSLTLLRAKGKEAIS 120
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; PRIORITY APPLICATION NUMBER: 10/658, 834
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; PRIORITY FILING DATE: 2003-03-21
; PRIORITY APPLICATION NUMBER: 60/409, 898
; PRIORITY FILING DATE: 2002-09-09
; NUMBER OF SEQ ID NOS: 1306
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 959
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; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-176-830-959

RESULT 6
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Best Local Similarity 99.4%; Pred. No. 2e-84; Matches 1; Mismatches 0; Indels 0; Gaps 0;
Matches 164; Conservative 1; Gaps 0;
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Number US-11-176-830-957
; FILE REFERENCE: 17109-012002 (922B)
; CURRENT APPLICATION NUMBER: US/11/176, 830
; PRIORITY FILING DATE: 2003-09-08
; PRIORITY APPLICATION NUMBER: 10/658, 834
; PRIORITY FILING DATE: 2005-07-06
; PRIORITY APPLICATION NUMBER: 60/457, 135
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; PRIORITY APPLICATION NUMBER: 60/409, 898
; PRIORITY FILING DATE: 2002-09-09
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; SEQ ID NO: 952
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; ORGANISM: Homo sapiens
; US-11-176-830-952

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Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 1; Mismatches 0; Indels 0; Gaps 0;
Matches 164; Conservative 1; Gaps 0;
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Number US-11-176-830-957
; FILE REFERENCE: 17109-012002 (922B)
; CURRENT APPLICATION NUMBER: US/11/176, 830
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; PRIORITY FILING DATE: 2003-03-21
; PRIORITY APPLICATION NUMBER: 60/409, 898
; PRIORITY FILING DATE: 2002-09-09
; NUMBER OF SEQ ID NOS: 1305
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 952
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-176-830-952

RESULT 8
Query Match 99.6%; Score 843; DB 7; Length 166;
Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 1; Mismatches 0; Indels 0; Gaps 0;
Matches 164; Conservative 1; Gaps 0;
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Number US-11-176-830-957
; FILE REFERENCE: 17109-012002 (922B)
; CURRENT APPLICATION NUMBER: US/11/176, 830
; PRIORITY FILING DATE: 2003-09-09
; PRIORITY APPLICATION NUMBER: 10/658, 834
; PRIORITY FILING DATE: 2005-07-06
; PRIORITY APPLICATION NUMBER: 60/457, 135
; PRIORITY FILING DATE: 2003-03-21
; PRIORITY APPLICATION NUMBER: 60/409, 898
; PRIORITY FILING DATE: 2002-09-09
; NUMBER OF SEQ ID NOS: 1306
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 955
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-176-830-967

Query Match 99.8%; Score 844; DB 7; Length 166;
Best Local Similarity 99.4%; Pred. No. 2e-84; Matches 1; Mismatches 0; Indels 0; Gaps 0;
Matches 164; Conservative 1; Gaps 0;
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Number US-11-176-830-957
; FILE REFERENCE: 17109-012002 (922B)
; CURRENT APPLICATION NUMBER: US/11/176, 830
; PRIORITY FILING DATE: 2003-09-09
; PRIORITY APPLICATION NUMBER: 10/658, 834
; PRIORITY FILING DATE: 2005-07-06
; PRIORITY APPLICATION NUMBER: 60/457, 135
; PRIORITY FILING DATE: 2003-03-21
; PRIORITY APPLICATION NUMBER: 60/409, 898
; PRIORITY FILING DATE: 2002-09-09
; NUMBER OF SEQ ID NOS: 1306
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 955
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-176-830-967

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; ORGANISM: Homo sapiens
; US-11-176-830-955
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; Db 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; QY 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; Db 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; QY 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; Db 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; ; ORGANISM: Homo sapiens
; US-11-176-830-958
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Sequence 958, Application US/111-76830
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Nu
; TITLE OF INVENTION: Acid Molecules and Related Applications
; FILE REFERENCE: 17109-012002 (922B)
; CURRENT APPLICATION NUMBER: US/111-176, 830
; CURRENT FILING DATE: 2005-07-06
; PRIOR APPLICATION NUMBER: 10/658, 834
; PRIOR FILING DATE: 2003-09-08
; PRIOR APPLICATION NUMBER: 60/457, 135
; PRIOR FILING DATE: 2003-03-21
; PRIOR APPLICATION NUMBER: 60/409, 898
; NUMBER OF SEQ ID NOS: 1306
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO: 958
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-11-176-830-958
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; Db 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; QY 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; Db 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; QY 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; Db 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; ; ORGANISM: Homo sapiens
; US-11-176-830-966
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; Db 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; QY 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; Db 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; QY 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; Db 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; ; ORGANISM: Homo sapiens
; RESULT 10
; US-11-176-830-966
; Sequence 966, Application US/111-76830
; Publication No. US20060020116A1
; GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; ; ORGANISM: Homo sapiens
; US-11-176-830-966
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Best Local Similarity 99.4%; Pred. No. 2.6e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; Db 1 APPRUCIDSVRVLRYLEAKAENITTCGCAEHCSLENENTVPDTKVNFKWAKMEVGQAA 60
; QY 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; Db 61 VEWVOGLALLSEAVLRLQGALLVNSQPWPPLQLHVDAVKGSLSTLRLGAQKEAIS 120
; QY 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; Db 121 PPDASAAPRTTADTRKLFRVYSNFGKLYTGEACRTGD 165
; ; ORGANISM: Homo sapiens
; RESULT 11
; US-11-181-091-34
; Query Match Score 99.6%; Score 843; DB 7; Length 166;
; Sequence 34, Application US/111-81091
; Publication No. US20060030046A1
; GENERAL INFORMATION:
; APPLICANT: Asada, Kiyozo
; APPLICANT: Uemori, Takashi
; APPLICANT: Kovama, Nobuto
; APPLICANT: Hashino, Kimikazu
; APPLICANT: Kato, Ikuoshin
; TITLE OF INVENTION: METHOD FOR GENE TRANSFER INTO TARGET CELLS WITH RETROVIRUS
; NUMBER OF SEQUENCES: 39
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: WEISER & ASSOCIATES
; STREET: 230 South Fifteenth Street, Suite 500
; CITY: Philadelphia
; STATE: PA
; COUNTRY: USA
; ZIP: 19102
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #11.0, Version #1.30
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/11/181, 091
; FILING DATE: 14-Jul-2005
; CLASSIFICATION: <Unknown>
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: US/09/775, 964
; FILING DATE: 20-Feb-2001
; APPLICATION NUMBER: US/09/366, 009
; FILING DATE: 02-Aug-1999
; APPLICATION NUMBER: 08/809, 156

FILING DATE: <Unknown>
APPLICATION NUMBER: JP 294382/1995
FILED DATE: 13-NOV-1995
APPLICATION NUMBER: JP 051847/1996
FILED DATE: 08-MAR-1996
ATTORNEY/AGENT INFORMATION:
NAME: Weiser, Gerard J.
REGISTRATION NUMBER: 19, 763
REFERENCE/DOCKET NUMBER: 977.6507P
TELECOMMUNICATION INFORMATION:
TELEPHONE: 215-875-8383
TELEFAX: 215-875-8394
INFORMATION FOR SEQ ID NO: 34:
SEQUENCE CHARACTERISTICS:
SEQUENCE LENGTH: 412 amino acids
TYPE: amino acid
STRANDEDNESS: <Unknown>
MOLECULE TYPE: peptide
SEQUENCE DESCRIPTION: SEQ ID NO: 34:
US-11-181-091-34

RESULT 12

Query Match Best Local Similarity 99.6%; Score 843; DB 7; Length 412;
Matches 164; Conservative 99.4%; Pred. No. 8, 8e-84; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENINIVDTKVNFYAWKMEVQQAA 60
Db 233 APPRLICDSRVLORYLLEAKEAENITGCAEHCSLENINIVDTKVNFYAWKMEVQQAA 292
Qy 61 VEVMOGALLSLSVRLRQALVNSQWEPQLQHVDAVSGLSLTTLRAKGKAIS 120
Db 293 VEVMOGALLSLSVRLRQALVNSQWEPQLQHVDAVSGLSLTTLRAKGKAIS 352
Qy 121 PPDASASAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 165
Db 353 PPDASASAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 397

RESULT 13

US-11-176-830-942
; Sequence 942, Application US/11176830
; Publication No. US20060202016A1
GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding N
; TITLE OF INVENTION: Acid Molecules and Related Applications
; FILE REFERENCE: 17109_012002 (922B)
; CURRENT FILING DATE: US/11-176, 830
; PRIOR APPLICATION NUMBER: 10/658, 834
; PRIOR FILING DATE: 2003-09-08
; PRIOR APPLICATION NUMBER: 60/457, 135
; PRIOR FILING DATE: 2003-03-21
; PRIOR APPLICATION NUMBER: 60/409, 898
; PRIOR FILING DATE: 2002-09-09
; NUMBER OF SEQ ID NOS: 1306
; SOFTWARE: FastSEQ for Windows Version 4.0
; SEQ ID NO 942
; LENGTH: 166
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Homo sapiens
US-11-176-830-942

RESULT 14

Query Match Best Local Similarity 99.5%; Score 842; DB 7; Length 166;
Matches 164; Conservative 99.4%; Pred. No. 3, 3e-84; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENINIVDTKVNFYAWKMEVQQAA 60
Db 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLENINIVDTKVNFYAWKMEVQQAA 60
Qy 61 VEVMOGALLSLSVRLRQALVNSQWEPQLQHVDAVSGLSLTTLRAKGKAIS 120
Db 61 VEVMOGALLSLSVRLRQALVNSQWEPQLQHVDAVSGLSLTTLRAKGKAIS 120
Qy 121 PPDASASAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 165
Db 121 PPDASASAPLRTTADTRKLFRVYSNPLRGKLYGEACRTGD 165

RESULT 15

US-11-176-830-948
; Sequence 948, Application US/11176830
; Publication No. US200602016A1
GENERAL INFORMATION:
; APPLICANT: Gantier, Rene
; APPLICANT: Guyon, Thierry
; APPLICANT: Drittanti, Lila
; APPLICANT: Vega, Manuel
; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding N
; TITLE OF INVENTION: Acid Molecules and Related Applications
; FILE REFERENCE: 17109_012002 (922B)
; CURRENT APPLICATION NUMBER: US/11/176, 830
; CURRENT FILING DATE: 2005-07-06

Query Match Best Local Similarity 99.6%; Score 843; DB 7; Length 444;
Matches 164; Conservative 99.4%; Pred. No. 9, 8e-84; Mismatches 0; Indels 0; Gaps 0;

; OTHER INFORMATION: Description of Artificial Sequence: Synthetic
; OTHER INFORMATION: Construct

US-11-029-003-16

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RESULT 17

RESULT 18

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RESULT 306

PRIOR APPLICATION NUMBER: 10/658,834

PRIOR FILING DATE: 2003-09-08

PRIOR APPLICATION NUMBER: 60/457,135

PRIOR FILING DATE: 2003-03-21

PRIOR APPLICATION NUMBER: 60/409,898

PRIOR FILING DATE: 2002-09-09

NUMBER OF SEQ ID NOS: 1306

SOFTWARE: FastSEQ for Windows Version 4.0

SEQ ID NO: 948

LENGTH: 166

TYPE: PRT

ORGANISM: Homo sapiens

US-11-176-830-948

Query Match Score 99.5%; Score 842; DB 7; Length 166;

Best Local Similarity 99.4%; Pred. No. 3.3e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLVEAENITTCGAEHCSENENTVDPDKVNPKWAKOMEVGQAA 60

Db 1 APPRLICDSRVLERYLVEAENITTCGAEHCSENENTVDPDKVNPKWAKOMEVGQAA 60

QY 61 VEWQGLALISEAVRGQALIVNSQWPBLQLANDKAVSGLSLTTLRAGQKEAIS 120

Db 61 VEWQGLALISEAVRGQALIVNSQWPBLQLANDKAVSGLSLTTLRAGQKEAIS 120

QY 121 PPDASAAPLRTTADFRKLFRVSNFLRGKLUYTGEACRTGD 165

Db 121 PPDASAAPLRTTADFRKLFRVSNFLRGKLUYTGEACRTGD 165

RESULT 15

US-11-176-830-951

; Sequence 951, Application US/11/176830

; Publication No. US20060020116A1

; GENERAL INFORMATION:

; APPLICANT: Gantier, Rene

; APPLICANT: Guyon, Thierry

; APPLICANT: Drittant, Lila

; TITLE OF INVENTION: Rational Evolution of Cytokines for Higher Stability, Encoding Nu-

; TITLE OF INVENTION: Acid Molecules and Related Applications

; FILE REFERENCE: 17109-012002 (922B)

; CURRENT APPLICATION NUMBER: US/11/176,830

; CURRENT FILING DATE: 2005-07-06

; PRIOR APPLICATION NUMBER: 10/658,834

; PRIOR FILING DATE: 2003-09-08

; PRIOR APPLICATION NUMBER: 60/457,135

; PRIOR FILING DATE: 2003-03-21

; PRIOR APPLICATION NUMBER: 60/409,898

; PRIOR FILING DATE: 2002-09-09

; NUMBER OF SEQ ID NOS: 1306

; SOFTWARE: FastSEQ for Windows Version 4.0

; SEQ ID NO: 951

; LENGTH: 166

; TYPE: PRT

; ORGANISM: Homo sapiens

US-11-176-830-951

Query Match Score 99.5%; Score 842; DB 7; Length 166;

Best Local Similarity 99.4%; Pred. No. 3.3e-84; Matches 164; Conservative 1; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLVEAENITTCGAEHCSENENTVDPDKVNPKWAKOMEVGQAA 60

Db 1 APPRLICDSRVLERYLVEAENITTCGAEHCSENENTVDPDKVNPKWAKOMEVGQAA 60

QY 61 VEWQGLALISEAVRGQALIVNSQWPBLQLANDKAVSGLSLTTLRAGQKEAIS 120

Db 61 VEWQGLALISEAVRGQALIVNSQWPBLQLANDKAVSGLSLTTLRAGQKEAIS 120

QY 121 PPDASAAPLRTTADFRKLFRVSNFLRGKLUYTGEACRTGD 165

Db 121 PPDASAAPLRTTADFRKLFRVSNFLRGKLUYTGEACRTGD 165

Search completed: February 28, 2006, 15:42:46

Job time : 18 secs

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run on: March 1, 2006, 10:19:01 ; Search time 47 Seconds
 (without alignments)
 sequence: (without alignments)
 scoring table: BLOSUM62
 Gapop 10.0 , Gapext 0.5

searched: 572060 seqs, 82675679 residues

total number of hits satisfying chosen parameters: 572060

maximum DB seq length: 0
 maximum DB seq length: 200000000
 listing first 45 summaries

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- 1: /cgn2_6_ptcodata/1/iaa/5_COMB.pep:*
- 2: /cgn2_6_ptcodata/1/iaa/6_COMB.pep:*
- 3: /cgn2_6_ptcodata/1/iaa/PCTUS_COMB.pep:*
- 4: /cgn2_6_ptcodata/1/iaa/RB_COMB.pep:*
- 5: /cgn2_6_ptcodata/1/iaa/backfile1.pep:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

result No.	Score	Query Length	DB ID	Description
1	846	100.0	165	2 US-09-604-871-1
2	846	100.0	165	2 US-09-604-938-1
3	846	100.0	165	2 US-09-830-967-1
4	846	100.0	165	2 US-10-241-356-1
5	846	100.0	166	1 US-08-318-193-70
6	846	100.0	166	2 US-09-604-871-2
7	846	100.0	166	2 US-09-604-938-2
8	846	100.0	166	2 US-09-462-941-2
9	846	100.0	166	2 US-10-360-101-227
10	846	100.0	166	2 US-10-241-356-2
11	846	100.0	166	4 PCT-US94-04361-37
12	846	100.0	193	1 US-07-903-220-1
13	846	100.0	193	1 US-08-353-918-34
14	846	100.0	193	1 US-08-883-795A-34
15	846	100.0	193	2 US-09-552-265B-4
16	846	100.0	193	2 US-09-813-775A-4
17	846	100.0	193	2 US-09-856-796B-4
18	846	100.0	435	2 US-09-932-812A-22
19	846	100.0	436	2 US-09-932-812A-18
20	846	100.0	437	2 US-09-932-812A-20
21	843	99.6	165	2 US-09-554-451-8
22	843	99.6	412	2 US-09-360-009-34
23	843	99.6	412	2 US-08-808-156B-34
24	843	99.6	412	2 US-09-775-964-34
25	838	99.1	193	2 US-09-552-265B-2
26	838	99.1	193	2 US-09-813-775C-2
27	834	98.6	193	2 US-09-552-265B-5

RESULT 1

; Sequence 1, Application US/09604871

; GENERAL INFORMATION:

; APPLICANT: Hilger, Bernd

; APPLICANT: Joesel, Hans-Peter

; TITLE OF INVENTION: ERYTHROPOIETIN CONJUGATES

; FILE REFERENCE: 1098 nonprovisional

; CURRENT APPLICATION NUMBER: US/09/604,871

; CURRENT FILING DATE: 2000-06-28

; PRIOR APPLICATION NUMBER: 60/151,454

; PRIOR FILING DATE: 1999-08-30

; PRIOR APPLICATION NUMBER: 60/147,452

; PRIOR FILING DATE: 1999-08-05

; PRIOR APPLICATION NUMBER: 60/142,243

; PRIOR FILING DATE: 1999-07-02

; NUMBER OF SEQ ID NOS: 3

; SOFTWARE: Patent In Ver. 2.1

; SEQ ID NO 1

; LENGTH: 165

; TYPE: PRT

; ORGANISM: Homo sapiens

US-09-604-871-1

Query Match 100.0%; Score 846; DB 2 Best Local Similarity 100.0%; Pre. No. 1.4e-9 Matches 165; Conservative 0; Mismatches 0

QY 1 APRILICDSRVLEERYLVEAKENNTTGCACHEHCSLNFE

Db 1 APRILICDSRVLEERYLVEAKENNTTGCACHEHCSLNFE

QY 61 VEWVOGALLISBEAVLRGQALIVNSQWPWEPQLQHVDK

Db 61 VEWVOGALLISBEAVLRGQALIVNSQWPWEPQLQHVDK

QY 121 PPDAASAPARTITADTRKLFRVSNFLAGKLKLYT

Db 121 PPDAASAPARTITADTRKLFRVSNFLAGKLKLYT

RESULT 2

; Sequence 1, Application US/09604938

; Patent No. 6583272

; GENERAL INFORMATION:

; APPLICANT: Ballon, Pascal

; TITLE OF INVENTION: ERYTHROPOIETIN CONJUGATES

FILE REFERENCE: 1097 nonprovisional
; CURRENT APPLICATION NUMBER: US/09/604,938
; CURRENT FILING DATE: 2000-06-27
; PRIORITY APPLICATION NUMBER: 60/166,151
; PRIORITY FILING DATE: 1999-11-17
; PRIORITY APPLICATION NUMBER: 60/151,548
; PRIORITY FILING DATE: 1999-08-13
; PRIORITY APPLICATION NUMBER: 60/150,225
; PRIORITY APPLICATION NUMBER: 60/142,254
; PRIORITY FILING DATE: 1999-07-02
; NUMBER OF SEQ ID NOS: 3
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-604-938-1

Query Match Score 846; DB 2; Length 165;
; Best Local Similarity 100.0%; Pred. No. 1.4e-99;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; Db 61 VEWQGLALLSEAVLRGQALLNSQWPBPLQLHVDKAVSGLSLTTLRALGAQEKAIS 120
; QY 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165
; Db 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165

RESULT 3
; US-09-830-967-1
; Sequence 1, Application US/09830967
; Patent No. 6777205
; GENERAL INFORMATION:
; APPLICANT: Sternbeld Biotechnology No. 6777205th America, Inc.
; APPLICANT: Carcagno, Carlos Miguel
; APPLICANT: Cribciolo, Marcelo
; APPLICANT: Melo, Carlos
; APPLICANT: Vidal, Juan Alejandro
; TITLE OF INVENTION: Host Cells Expressing Recombinant Human Erythropoietin
; FILE REFERENCE: 1909-020002
; CURRENT FILING DATE: 1999-11-08
; PRIOR APPLICATION NUMBER: AR 99-01-00679
; PRIOR FILING DATE: 1999-02-23
; PRIOR APPLICATION NUMBER: AR 98-01-05609
; PRIOR FILING DATE: 1998-11-06
; NUMBER OF SEQ ID NOS: 5
; SOFTWARE: PatentIn Ver. 2.0
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-09-830-967-1

Query Match Score 846; DB 2; Length 165;
; Best Local Similarity 100.0%; Pred. No. 1.4e-99;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; Db 61 VEWQGLALLSEAVLRGQALLNSQWPBPLQLHVDKAVSGLSLTTLRALGAQEKAIS 120
; QY 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165
; Db 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165

RESULT 4
; US-10-241-356-1
; Sequence 1, Application US/10241356
; Patent No. 6930086
; GENERAL INFORMATION:
; APPLICANT: TISCHER, WILHELM
; TITLE OF INVENTION: DIGLYCOSYLATED ERYTHROPOETIN
; FILE REFERENCE: 20971
; CURRENT APPLICATION NUMBER: US/10/241,356
; CURRENT FILING DATE: 2002-09-11
; PRIORITY APPLICATION NUMBER: EP 01122555.4
; PRIORITY FILING DATE: 2001-09-25
; NUMBER OF SEQ ID NOS: 2
; SOFTWARE: PatentIn Ver. 2.1
; SEQ ID NO 1
; LENGTH: 165
; TYPE: PRT
; ORGANISM: Homo sapiens
; US-10-241-356-1

Query Match Score 846; DB 2; Length 165;
; Best Local Similarity 100.0%; Pred. No. 1.4e-99;
; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
; QY 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; 1 APPRLICDSRVLERYLLEAKEAENITGCAEHCSLNENITVDPDKVNFKWAKMEVGQAA 60
; Db 61 VEWQGLALLSEAVLRGQALLNSQWPBPLQLHVDKAVSGLSLTTLRALGAQEKAIS 120
; QY 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165
; Db 121 PPDAASAPLRTTADTRKLFRVYSNPLRGKLUYTGEACRTGD 165

RESULT 5
; US-08-318-193-70
; Sequence 70, Application US/08318193
; Patent No. 5641663
; GENERAL INFORMATION:
; APPLICANT: GARVIN, Robert T.
; APPLICANT: MALEK, Lawrence T.
; TITLE OF INVENTION: AN EXPRESSION SYSTEM FOR THE SECRETION
; TITLE OF INVENTION: OF BIOACTIVE HUMAN GRANULOCYTE MACROPHAGE COLONY
; TITLE OF INVENTION: STIMULATING FACTOR (GM-CSF) AND OTHER HETEROLOGOUS
; TITLE OF INVENTION: PROTEINS FROM STREPTOMYCES
; NUMBER OF SEQUENCES: 91
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Foley & Lardner
; STREET: 1800 Diagonal Road, Suite 500
; CITY: Alexandria
; STATE: Virginia
; COUNTRY: USA
; ZIP: 22313-0239
; COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: PatentIn Release #1.0, Version #1.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/08/318,193
; FILING DATE:
; CLASSIFICATION: 435
; PRIORITY APPLICATION DATA:
; APPLICATION NUMBER: US/07/935,314

FILING DATE: APPLICATION NUMBER: US 07/224,568
 ATTORNEY/AGENT INFORMATION:
 NAME: BENT, Stephen A.
 REGISTRATION NUMBER: 29,768
 REFERENCE/DOCKET NUMBER: 10740/116 CACO
 TELECOMMUNICATION INFORMATION:
 TELEPHONE: (703) 833-9300
 TELEFAX: 899149
 INFORMATION FOR SEQ ID NO: 70:
 SEQUENCE CHARACTERISTICS:
 LENGTH: 166 amino acids
 TYPE: amino acid
 TOPOLOGY: linear
 MOLECULE TYPE: protein
 ;US-08-318-193-70

Query Match 100.0%; Score 846; DB 1; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.4e-99;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60
 DB 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60
 QY 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120
 DB 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120

QY 121 PPDAAASAPLRTITADTPRKLFPRVVSNPLRGKLUYTGEACRTGD 165
 DB 121 PPDAAASAPLRTITADTPRKLFPRVVSNPLRGKLUYTGEACRTGD 165

RESULT 6
 US-08-604-871-2
 Sequence 2, Application US/09604871
 ; Patent No. 6340742
 ; GENERAL INFORMATION:
 ; APPLICANT: Burg, Josef
 ; APPLICANT: Hilger, Bernd
 ; APPLICANT: Josel, Hans-Peter
 TITLE OF INVENTION: ERYTHROPOETIN CONJUGATES
 FILE REFERENCE: 1098 nonprovisional
 CURRENT APPLICATION NUMBER: US/08/ 604, 871
 CURRENT FILING DATE: 2000-06-28
 PRIOR APPLICATION NUMBER: 60/151, 454
 PRIOR FILING DATE: 1999-08-30
 PRIOR APPLICATION NUMBER: 60/147, 452
 PRIOR FILING DATE: 1999-08-05
 PRIOR APPLICATION NUMBER: 60/142, 243
 PRIOR FILING DATE: 1999-07-02
 NUMBER OF SEQ ID NOS: 3
 SOFTWARE: PatentIn Ver. 2.1
 SEQ ID NO 2
 LENGTH: 166
 TYPE: PRI
 ORGANISM: Homo sapiens
 ;US-08-604-871-2

Query Match 100.0%; Score 846; DB 2; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.4e-99;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60
 DB 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60

QY 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120
 DB 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120

QY 121 PPDAAASAPLRTITADTPRKLFPRVVSNPLRGKLUYTGEACRTGD 165
 DB 121 PPDAAASAPLRTITADTPRKLFPRVVSNPLRGKLUYTGEACRTGD 165

RESULT 8
 US-09-462-941-2
 Sequence 2, Application US/09462941
 ; Patent No. 6608183
 ; GENERAL INFORMATION:
 ; APPLICANT: Cox III, George N
 ; APPLICANT: Bolder Biotechnology, Inc.
 TITLE OF INVENTION: Derivatives of Growth Hormone and Related Proteins
 FILE REFERENCE: 4152-1-UPS
 CURRENT APPLICATION NUMBER: US/09/462, 941
 CURRENT FILING DATE: 2000-01-14
 PRIOR APPLICATION NUMBER: 60/052, 516
 PRIOR FILING DATE: 1997-07-14
 NUMBER OF SEQ ID NOS: 41
 SOFTWARE: PatentIn Ver. 2.0
 SEQ ID NO 2
 LENGTH: 166
 TYPE: PRT
 ORGANISM: Homo sapiens
 ;US-09-462-941-2

Query Match 100.0%; Score 846; DB 2; Length 166;
 Best Local Similarity 100.0%; Pred. No. 1.4e-99;
 Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60
 DB 1 APPRLICDSRVLERYVLEAKENAINTGCAHCISLNENITVPDTKVNFYAWKRMEVGQA 60

QY 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120
 DB 61 VEWQGALLSEAVRGQALLVNSQWEPQLHYDKAVSGRSLLTLLRIGAQUEAIS 120

QY 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
; 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
Db 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
; 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
QY 121 PPDASAAPLRTTADTRKLFRYVSNFLRGKLUYGEACRTGD 165
Db 121 PPDASAAPLRTTADTRKLFRYVSNFLRGKLUYGEACRTGD 165
; RESULT 9
US-10-360-101-227
; Sequence 227, Application US/10360101
; Patent No. 6861236
; GENERAL INFORMATION:
; APPLICANT: Moll, Gert N.
; INVENTION: Leenhouts, Cornelis J.
; TITLE OF INVENTION: Export modification of (poly)peptide in the lantibiotic way
; FILE REFERENCE: 2183-5673
; CURRENT APPLICATION NUMBER: US/10/360,101
; CURRENT FILING DATE: 2003-07-07
; PRIOR APPLICATION NUMBER: EP 02077060.8
; PRIOR FILING DATE: 2002-05-24
; NUMBER OF SEQ ID NOS: 309
; SOFTWARE: Patentin version 3.1
; SEQ ID NO 227
; LENGTH: 166
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE:
; OTHER INFORMATION: sequence of erythropoietin
; US-10-360-101-227

Query Match 100.0%; Score 846; DB 2; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.4e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
Db 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
; 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
QY 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
Db 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
; 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
QY 121 PPDASAAPLRTTADTRKLFRYVSNFLRGKLUYGEACRTGD 165
Db 121 PPDASAAPLRTTADTRKLFRYVSNFLRGKLUYGEACRTGD 165
; RESULT 11
PCT-US94-04361-37
; Sequence 37, Application PC/TUS9404361
; GENERAL INFORMATION:
; APPLICANT: Brigham and Women's Hospital
; APPLICANT: 75 Francis Street
; APPLICANT: Boston, MA 02115
; APPLICANT: Burn, H. Franklin
; APPLICANT: Wen, Danyi
; APPLICANT: Showers, Mark O.
; TITLE OF INVENTION: Erythropoietin Muteins With Enhanced
; TITLE OF INVENTION: Activity
; NUMBER OF SEQIDNOS: 59
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Serne, Kessler, Goldstein & Fox
; STREET: 1100 New York Avenue, Suite 600
; CITY: Washington
; STATE: D.C.
; COUNTRY: U.S.A.
; ZIP: 20005-3334
; COMPUTER READABLE FORM:
; MEDIUM TYPE:
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentin Release #1.0, Version #11.25
; CURRENT APPLICATION DATA:
; APPLICATION NUMBER: PCT/US94/04361
; FILING DATE: Herewith
; CLASSIFICATION:
; PRIOR APPLICATION DATA:
; APPLICATION NUMBER: 08/049, 802
; FILING DATE: 21-APR-1993
; ATTORNEY/AGENT INFORMATION:
; NAME: Climalia, Michele A.
; REGISTRATION NUMBER: 33, 851
; REFERENCE/DOCKET NUMBER: 0627.336PC01
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 371-2000
; TELEFAX: (202) 371-2540
; INFORMATION FOR SEQ ID NO: 37:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 166 amino acids
; TYPE: amino acid
; TOPOLOGY: both
; PCT-US94-04361-37

Query Match 100.0%; Score 846; DB 4; Length 166;
Best Local Similarity 100.0%; Pred. No. 1.4e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
Db 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
; 1 APPRLICDSRVLEERYLLEAKEAENITGCAEHCSLENINITPDTKVNLYAWKRMEVGQA 60
QY 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120
; 61 VEWMOGLALLSEAVLRGOALLVNSQPWEPLQLHVDKAVSGRSLLTIRALGAOKEALS 120

RESULT 12
US-07-903-220-1
; Sequence 1, Application US/07903220
; Patent No. 5322337
; GENERAL INFORMATION:
; APPLICANT: Hewick, Rodney M.
; TITLE OF INVENTION: METHOD FOR THE PURIFICATION OF
; ERYTHROPOIETIN AND ERYTHROPOIETIN COMPOSITION
; NUMBER OF SEQUENCES: 1
; CORRESPONDENCE ADDRESS:
; ADDRESSEE: Paul H. Heller
; STREET: Kenyon & Kenyon, One Broadway
; CITY: New York
; STATE: New York
; COUNTRY: USA
; ZIP: 10004

COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentin Release #1.0, Version #1.25

CURRENT APPLICATION DATA:
; APPLICATION NUMBER: US/07/903, 220
; FILING DATE: 19920731
; CLASSIFICATION: 530

ATTORNEY/AGENT INFORMATION:
; NAME: Brown, Scott A.
; REFERENCE/DOCKET NUMBER: 32, 724
; TELECOMMUNICATION INFORMATION:
; TELEPHONE: (202) 429-1776
; TELEFAX: (202) 429-0796
; INFORMATION FOR SEQ ID NO: 1:
; SEQUENCE CHARACTERISTICS:
; LENGTH: 193 amino acids
; TYPE: AMINO ACID
; TOPLOGY: Linear
; MOLECULE TYPE: protein
; HYPOTHETICAL: NO
; ORIGINAL SOURCE:
; ORGANISM: Homo sapiens
; US-07-903-220-1

Query Match 100.0%; Score 846; DB 1; Length 193;
Best Local Similarity 100.0%; Pred. No. 1.8e-99;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

Qy 1 APPRLICDSVLERLVEAKAENTTGCACRHCSLENITVPDTKVNPKYAKRMEVGQQA 60
Db 28 APPRLICDSVLERLVEAKAENTTGCACRHCSLENITVPDTKVNPKYAKRMEVGQQA 87

Qy 61 VEVWQGLALISEAVIRGQALVNSSQPWEPLQLHDKAVSGLSLTLLRAGAQKAIIS 120
Db 88 VEVWQGLALISEAVIRGQALVNSSQPWEPLQLHDKAVSGLSLTLLRAGAQKAIIS 147

Qy 121 PPDASASAAPLTTTADTPRKFLFRVSNFLRGKLKLYTGEACRTGD 165
Db 148 PPDASASAAPLTTTADTPRKFLFRVSNFLRGKLKLYTGEACRTGD 192

RESULT 13
US-08-388-918-34
; sequence 34, Application US/08388918
; Patent No. 588774
; GENERAL INFORMATION:
; APPLICANT: Delcuve, Genevieve

GENERAL INFORMATION:
; APPLICANT: Delcuve, Genevieve
; TITLE OF INVENTION: Recombinant DNA Molecules and Expression
; TITLE OF INVENTION: Vectors for Tissue Plasminogen Activator
; NUMBER OF SEQUENCES: 39

CORRESPONDENCE ADDRESS:
; ADDRESSEE: BERESIN & PARR
; STREET: 40 King Street West
; CITY: Toronto
; STATE: Ontario
; COUNTRY: Canada
; ZIP: M5H 3Y2

COMPUTER READABLE FORM:
; MEDIUM TYPE: Floppy disk
; COMPUTER: IBM PC compatible
; OPERATING SYSTEM: PC-DOS/MS-DOS
; SOFTWARE: Patentin Release #1.0, Version #1.25

RESULT 14
US-08-883-795A-34
; Sequence 34, Application US/08883795A
; Patent No. 5885607
; GENERAL INFORMATION:
; APPLICANT: Awang, Gregor
; TITLE OF INVENTION: Recombinant DNA Molecules and Expression
; TITLE OF INVENTION: Vectors for Tissue Plasminogen Activator
; NUMBER OF SEQUENCES: 39

CORRESPONDENCE ADDRESS:
; ADDRESSEE: BERESIN & PARR
; STREET: 40 King Street West
; CITY: Toronto
; STATE: Ontario
; COUNTRY: Canada
; ZIP: M5H 3Y2

RESULT 15
US-08-388-918-34
; sequence 34, Application US/08388918
; Patent No. 588774
; GENERAL INFORMATION:
; APPLICANT: Delcuve, Genevieve

CURRENT APPLICATION DATA:

APPLICATION NUMBER: US/08/883,795A

FILING DATE: 27-JUN-1997

CLASSIFICATION: 435

ATTORNEY/AGENT INFORMATION:

NAME: Gravelle, Micheline

REGISTRATION NUMBER: 40,261

REFERENCE/DOCKET NUMBER: 7841-062

TELECOMMUNICATION INFORMATION:

TELEPHONE: (416) 361-7311

TELEFAX: (416) 361-1398

INFORMATION FOR SEQ ID NO: 34:

SEQUENCE CHARACTERISTICS:

LENGTH: 193 amino acids

TYPE: amino acid

TOPOLOGY: linear

MOLECULE TYPE: protein

US-08-883-795A-34

Query Match 100.0%; Score 846; DB 1; Length 193;

Best Local Similarity 100.0%; Pred. No. 1.8e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 60

Db 28 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 87

QY 61 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 120

Db 88 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 147

QY 121 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 165

Db 148 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 192

RESULT 15
US-09-552-265B-4

Query Match 100.0%; Score 846; DB 1; Length 193;

Best Local Similarity 100.0%; Pred. No. 1.8e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 60

Db 28 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 87

QY 61 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 120

Db 88 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 147

QY 121 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 165

Db 148 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 192

RESULT 16
US-09-552-265B-4Query Match 100.0%; Score 846; DB 2; Length 193;
Title of Invention: polypeptides and nucleic acids encoding the same
FILE REFERENCE: GENENT.057CPI
CURRENT APPLICATION NUMBER: US 09/552,265B
CURRENT FILING DATE: 2000-04-19
PRIORITY APPLICATION NUMBER: US 09/307307
PRIOR FILING DATE: 1999-05-17
NUMBER OF SEQ ID NOS: 49
SOFTWARE: FastSBQ for Windows Version 4.0
SEQ ID NO 4
LENGTH: 193
TYPE: PRT
ORGANISM: Homo sapiens

US-09-552-265B-4

Query Match 100.0%; Score 846; DB 2; Length 193;

Best Local Similarity 100.0%; Pred. No. 1.8e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 60

Db 28 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 87

QY 61 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 120

Db 88 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 147

QY 121 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 165

Db 148 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 192

RESULT 17
US-09-856-796B-4Query Match 100.0%; Score 846; DB 2; Length 193;
Title of Invention: NF-kB ACTIVATION INHIBITORS, AND THEIR PHARMACEUTICAL
FILE REFERENCE: US898CMN
CURRENT APPLICATION NUMBER: US/09/856,796B
CURRENT FILING DATE: 2001-05-07
PRIORITY APPLICATION NUMBER: PCM/FRS9/02197
PRIOR FILING DATE: 1998-11-24
PRIORITY APPLICATION NUMBER: FR 98/14858
NUMBER OF SEQ ID NOS: 4

QY 1 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 60

Db 28 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 87

QY 61 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 120

Db 88 VEWVOGLALISSEAVLRGQALLNSQWPFLQLVHDVAVGSLSLTTLRAGQKEAIS 147

QY 121 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 165

Db 148 PPDASAASAPLRTTADTRKLFRTVSNPLRGKLYTGEACTGD 192

Query Match 100.0%; Score 846; DB 2; Length 193;

Best Local Similarity 100.0%; Pred. No. 1.8e-99; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;

QY 1 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 60

Db 67 APPRLICDSRVLERYLLEAKENAENITGCAEHCSLNENITVDPDKVNFTWAKRMEVGQQA 87

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Db 28 APPRICDSRVRLERYLLEAKEAENITTCGCAEHCSLNENITVDPDKVNPFYAWKMEVGQAA 87 ; ORGANISM: Artificial Sequence
Qy 61 VEWQGLALLSEAVLRGQALLVNSSQWPFLQLHDKAVSGLSLTTLRGAQEKAIS 120 ; FEATURE: FEATUR
; E: OTHER INFORMATION: HUEPO-L-vPC gamma2 with a 27-amino acid leader peptide
Db 88 VEWQGLALLSEAVLRGQALLVNSSQWPFLQLHDKAVSGLSLTTLRGAQEKAIS 147 ; OTHER INFORMATION: (Figure 2
; OTHER INFORMATION: A)
Qy 121 PPDRASAAPLRTTADTRKLFRVYSNFRGKUKLYGEACRGD 165 ; US-09-932-812A-18
Db 148 PPDRASAAPLRTTADTRKLFRVYSNFRGKUKLYGEACRGD 192 ; Sequence 22, Application US/09932812A
; Patent No. 6900292
; GENERAL INFORMATION:
; APPLICANT: Sun, Lee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: FC fusion proteins of human erythropoietin with
; TITLE OF INVENTION: increased biological
; TITLE OF INVENTION: activities
; FILE REFERENCE: 02SUN2001
; CURRENT APPLICATION NUMBER: US/09/932,812A
; CURRENT FILING DATE: 2001-08-17
; NUMBER OF SEQ ID NOS: 28
; SEQ ID NO 22
; LENGTH: 435
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE: OTHER INFORMATION: HuEPO-L-vPC gamma1 with a 27-amino acid leader peptide
; OTHER INFORMATION: (Figure 2C
; OTHER INFORMATION: )
US-09-932-812A-22 ; US-09-932-812A-18
Query Match Score 846; DB 2; Length 435; ; Query Match Score 846; DB 2; Length 436;
Best Local Similarity 100.0%; Pred. No. 6.4e-99; ; Best Local Similarity 100.0%; pred. No. 6.5e-99;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0; ; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 APPRICDSRVRLERYLLEAKEAENITTCGCAEHCSLNENITVDPDKVNPFYAWKMEVGQAA 60 ; US-09-932-812A-20
Db 28 APPRICDSRVRLERYLLEAKEAENITTCGCAEHCSLNENITVDPDKVNPFYAWKMEVGQAA 87 ; Sequence 20, Application US/09932812A
; Patent No. 6900292
; GENERAL INFORMATION:
; APPLICANT: Sun, Lee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: FC fusion proteins of human erythropoietin with
; TITLE OF INVENTION: increased biological
; TITLE OF INVENTION: activities
; FILE REFERENCE: 02SUN2001
; CURRENT APPLICATION NUMBER: US/09/932,812A
; CURRENT FILING DATE: 2001-08-17
; NUMBER OF SEQ ID NOS: 28
; SEQ ID NO 20
; LENGTH: 437
; TYPE: PRT
; ORGANISM: Artificial Sequence
; FEATURE: OTHER INFORMATION: HuEPO-L-vPC gamma2 with a 27-amino acid leader peptide
; OTHER INFORMATION: (Figure 2B
; OTHER INFORMATION: )
US-09-932-812A-20 ; US-09-932-812A-20
Query Match Score 846; DB 2; Length 437; ; Query Match Score 846; DB 2; Length 437;
Best Local Similarity 100.0%; Pred. No. 6.5e-99; ; Best Local Similarity 100.0%; pred. No. 6.5e-99;
Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0; ; Matches 165; Conservative 0; Mismatches 0; Indels 0; Gaps 0;
Qy 1 APPRICDSRVRLERYLLEAKEAENITTCGCAEHCSLNENITVDPDKVNPFYAWKMEVGQAA 60 ; US-09-932-812A-18
Db 28 APPRICDSRVRLERYLLEAKEAENITTCGCAEHCSLNENITVDPDKVNPFYAWKMEVGQAA 87 ; Sequence 18, Application US/09932812A
; Patent No. 6900292
; GENERAL INFORMATION:
; APPLICANT: Sun, Lee-Hwei K
; APPLICANT: Sun, Bill N
; APPLICANT: Sun, Cecily R
; TITLE OF INVENTION: FC fusion proteins of human erythropoietin with
; TITLE OF INVENTION: increased biological
; TITLE OF INVENTION: activities
; FILE REFERENCE: 02SUN2001
; CURRENT APPLICATION NUMBER: US/09/932,812A
; CURRENT FILING DATE: 2001-08-17
; NUMBER OF SEQ ID NOS: 28
; SOFTWARE: PatentIn version 3.1
; SEQ ID NO 18
; LENGTH: 436
; TYPE: PRT

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Search completed: March 1, 2006, 10:20:08
Job time : 48 secs

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